Long-term and persistent vocal plasticity in adult bats

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Bats exhibit a diverse and complex vocabulary of social communication calls some of which are believed to be learned during development. This ability to produce learned, species-specific vocalizations – a rare trait in the animal kingdom – requires a high-degree of vocal plasticity. Bats live extremely long lives in highly complex and dynamic social environments, which suggests that they might also retain a high degree of vocal plasticity in adulthood, much as humans do. Here, we report persistent vocal plasticity in adult bats (Rousettus aegyptiacus) following exposure to broad-band, acoustic perturbation. Our results show that adult bats can not only modify distinct parameters of their vocalizations, but that these changes persist even after noise cessation – in some cases lasting several weeks or months. Combined, these findings underscore the potential importance of bats as a model organism for studies of vocal plasticity, including in adulthood.
Humans are renowned for the ability to learn their vocalizations and maintain vocal plasticity throughout life. This ability allows them to modify and adjust properties of their vocal repertoire in response to auditory inputs from their surroundings. Amongst animals (particularly mammals) very few species share this capacity, resulting in a sparsity of relevant model organisms for neurobiological studies of this trait. To date, nearly all detailed neurobiological investigations have been conducted in birds, with most focusing on plasticity in juveniles but see Tumer and Brainard. These studies have yielded invaluable insight and revealed many of the computations that might support the bird’s capacity of song imitation as well as its potential parallel to human speech. At the same time, the field has also faced translational challenges due to evolutionary divergence between avian and mammalian brains (e.g. the nuclear organization of the avian pallium vs. the six-layered organization of mammalian cortex). Thus, the absence of a tractable mammalian model system for studying mechanisms of vocal plasticity, particularly in adulthood, represents a major and long-standing gap in the field.

Recently, bats have been proposed as a potential mammalian model system for bridging this gap owing to their unique specialization in acoustic signaling. In addition to their extraordinary echolocation capability, bats also possess a rich and diverse repertoire of social communication calls that are used primarily for social interactions. Importantly, some of the bats’ social vocalizations are believed to be learned during development by attending to auditory signaling produced by conspecifics in their surroundings. This rare ability is posited to be an extreme form of long-term and persistent vocal plasticity in a developing mammal.

To what extent is bat vocal plasticity retained in adulthood? The ethology of a bat’s life suggests that maintaining the capacity for vocal plasticity beyond development could be extremely advantageous. Bats are amongst the longest living mammals (comparing lifespan relative to body size) and spend much of their lives in socially complex and highly dynamic environments where effective acoustic interactions need to occur under noisy conditions. For example, Egyptian fruit bats (Rousettus aegyptiacus) live in large cave dwellings with many thousands of individuals where ambient noise levels can reach up to 100 dB SPL (Y. Yovel, personal communication). Under such varying and complex acoustic conditions, bats would clearly benefit from retaining a high degree of vocal plasticity as part of an adaptive, robust and effective vocal communication system. This notion, however, remains largely unexplored, especially in response to structured acoustic perturbation under controlled laboratory conditions (but see Boughman et al.).

Here, we set out to examine the extent of vocal plasticity in adult Egyptian fruit bats for two major reasons. First, recent studies have demonstrated the crucial role of auditory feedback during development in shaping their social vocalizations, underscoring the potential validity of this species as a mammalian model organism for vocal plasticity and learning. Second, a suite of novel methodologies has been developed and customized for adult Egyptian fruit bats, enabling neurophysiological measurements during uninhibited social interactions and during free flight. Yet, two major questions must first be addressed in order to support the use of adult Egyptian fruit bats for studies of vocal plasticity under laboratory conditions. First, does this species retain a capacity for vocal plasticity in adulthood, and if so, can it be engaged and studied under laboratory conditions? Addressing these two major questions was the goal of the present study where we leveraged the natural behavior of Egyptian fruit bats to vocally communicate even under acoustically noisy conditions.

Acoustic noise can mask communication signals, thereby disrupting information transfer and leading to errors in detection or interpretation of the signals by the receiver. It has been shown that most animals—from insects, amphibians, and birds to mammals—modify their acoustic signals (amplitude, duration and spectral composition) to maintain the signal-to-noise ratio of their vocalizations in the presence of noise, thereby increasing the probability of successful communication. Therefore, we designed an ethological behavioral paradigm to investigate how bats adapt the acoustic parameters of their vocalizations to the spectral features of their surroundings under controlled laboratory conditions (Fig. 1). Importantly, the focus of this study was to specifically assess whether these adaptive changes would persist after cessation of the acoustic perturbation (i.e., in the complete absence of noise), which suggests long-term and persistent vocal plasticity.

In its social environment, the Egyptian fruit bat is a highly vocal bat, and its hearing sensitivity and call frequency distribution are well characterized. By specifically tailoring spectral shapes of a continuous, broadband background noise, we were able to target the bats’ most sensitive frequency range or the most dominant call frequency range of their social calls (see Methods). We hypothesized that the bats’ communication calls should exhibit changes in their spectral features due to the influences of the presented acoustic perturbation. Figure 1 depicts a theoretical shift of call frequency correlated with spectral properties of the noise, where the mean magnitude spectrum of the bats’ calls before and after a theoretical shift is depicted in black and red (dashed line), respectively. When exposed to a noise with more energy in distinct frequencies, the bats can acoustically “escape” into lower or higher frequency bands to optimize their signal-to-noise ratio. Since the background noise was presented to the bats continuously (requiring them to communicate acoustically in noisier conditions), we hypothesized that the observed frequency shift in the calls might persist even after the noise was silenced—indicative of vocal plasticity. Hence, after noise cessation, we continuously monitored all social communication calls occurring during the bats’ natural interactions to assess the long-term effect of the acoustic interference.

We find that after noise cessation, adult bats modify distinct parameters of their vocalizations to minimize interference with the acoustic perturbation, and the observed shifts persist for up to several weeks or months after noise cessation. The obtained results demonstrate that this species retains the capacity for vocal plasticity in adulthood, underscoring its potential importance as a model organism for studies of vocal plasticity throughout stages of development.

**Results**

**Vocal plasticity in response to acoustic perturbation.** Bats rarely call in isolation and interact in very close proximity to each other such that individual identification of who is calling is unreliable. We thus recorded from six independent groups of bats. Two of the six groups exhibited a very low call rate of <80 calls per day during the baseline period prior to noise manipulation and were therefore excluded from the experiment and not subjected to the noise perturbation. The nature of vocal interaction in this species of bats is pairwise, and in agreement with previous reports we did not observe individual bats dominating the ‘conversation’. We most often observed multiple differently composed groups of bats engaging in vocal interactions, as reported previously (ref. 31).

We initially examined several distinct noise conditions: the first noise condition (noise #1, pink curve, Fig. 1c and Supplementary Fig. 1a) spanned much of the bats’ most sensitive hearing range.
and call spectrum (8–27 kHz). The second noise condition (noise #2, blue curve, Fig. 1c and Supplementary Fig. 1a), displayed a steeper slope and had a higher magnitude in the more dominant frequencies of the Egyptian fruit bat call spectrum (4–20 kHz). A third noise condition (noise #3, presented only to Group 1, Supplementary Fig. 1A) had the steepest spectral slope and energy mainly in the low frequency range (below 5 kHz). The last noise condition was only tested in one group and hence not included in the main analysis but is reported in Supplementary Fig. 1. Before noise presentation, all groups were acclimated for one week to the housing environment in which the acoustic perturbation took place. After 2 additional weeks of baseline recording, noise perturbation commenced. The acoustic perturbation consisted of low amplitude (45–50 dB, as used in prior studies in bats\textsuperscript{32,33}), continuous presentation of background noise over a period of 2 weeks. This amplitude level was chosen in order to elicit changes in vocalizations without disturbing the bats such that they would cease their vocal interactions. In total, we recorded 6348, 17115, 14153, and 10256 calls during the baseline period for Groups 1, 2, 3, and 4, respectively. The critical measurements of the experiment occurred following the cessation of the acoustic perturbation periods where we recorded the bats’ natural interactions (see examples in Supplementary Fig. 2).

The first two groups in the experiment consisted of six bats each that were recorded in a large enclosure (Fig. 1a; see Methods). The background noise was designed to contain more energy in the most sensitive hearing range (8–27 kHz) of the adult bat and tapered off toward the lower and higher frequencies (Fig. 1c and Supplementary Fig. 1A). We hypothesized that adult bats of this species might adapt their vocalizations to minimize interference with the presented noise perturbation. Furthermore, if adult bats do retain the capacity for vocal plasticity in adulthood, then this vocal adaptation should persist after cessation of the acoustic perturbation. Previous studies\textsuperscript{32,33} have shown that bats shift the centroid of their echolocation calls (amongst other call parameters) during noise presentation contingent upon background noise. Thus, initial examination of our hypothesis focused on the spectral centroid (the calls’ spectral center of mass)\textsuperscript{34} of the bats’ social vocalizations by comparing this measure before (baseline) and after presentation of each of the noise conditions. We hypothesized that any changes in the centroid should not result in strong reorganizations of the call structure, since only the energy distribution of the call is shifted within its own frequency range, allowing escape from the masking noise (see Fig. 1d).

During the week of silence after the presentation of noise #1 (pink curve in Fig. 1c) we recorded a total of 3442 calls for Group 1 and 10770 calls for Group 2. A summarization procedure into observation bins was applied on all calls (see Methods) and all further analysis was done on the summarized data sets. These
summarized calls were compared to all summarized calls recorded during baseline. We found that not only did the bats adapt their vocalizations by shifting their center of mass upwards but that this adaptation persisted in the complete absence of the noise (although this change was not significantly different from baseline when Bonferroni corrected for 11 multiple comparisons, see Methods; Fig. 1e, $1.39 \pm 0.623$ kHz for both groups, $t(8.66) = 2.23$ and $p = 0.028$, $n = 537$ bins, t-test through mixed linear model, see Methods). This finding was also not significant for both groups individually (Supplementary Fig. 1B,C; $0.471 \pm 0.75$ kHz, $z = 0.628$ and $p = 0.53$, $n = 139$ bins for Group 1, $1.69 \pm 0.811$ kHz, $z = 2.083$ and $p = 0.037$, $n = 398$ bins for Group 2, z-test through AR(1) model, see Methods).

The observed, moderate shift suggests that adult bats may retain a form of long-term plasticity for vocal production, which allows them to change the acoustic features of their vocalizations in response to changes in the acoustics of their environment. To further explore this hypothesis, we next examined whether the observed persistent shift depends on the spectral properties of the presented acoustic perturbation. To assess this possibility, both groups were subsequently exposed to noise #2, which differed in having a steeper spectral slope (blue curve in Fig. 1c and Supplementary Fig. 1A). We reasoned that because a larger portion of the bats’ most dominant call frequencies would now be perturbed, this manipulation would further drive the bats to increase their spectral center of mass to avoid the noise. During the week of silence after the presentation of noise #2, we recorded 5704 calls for Group 1 and 18272 calls for Group 2. Indeed, (following the same summarization procedure) we found that the mean centroid of the calls recorded after the cessation of noise #2 was further shifted upward compared to baseline by $3.118 \pm 0.545$ kHz for Group 1 and $2.795 \pm 0.515$ kHz for Group 2 ($z = 5.726$ and $p < 0.001$, $n = 171$ bins for Group 1, $z = 5.424$ and $p < 0.001$, $n = 505$ bins for Group 2, z-test through AR(1) model), as well as, when compared to noise #1 by $2.627 \pm 0.761$ kHz for Group 1 and $1.091 \pm 0.633$ kHz for Group 2 ($z = 3.452$ and $p < 0.001$, $n = 130$ bins for Group 1, $n = 415$ bins, $z = 1.723$ and $p = 0.085$ for Group 2, z-test through AR(1) model). The mean shift for both groups was $2.87 \pm 0.404$ kHz relative to baseline (Fig. 1e, $t(144) = 7.11$ and $p < 0.001$, $n = 676$ bins, t-test through mixed linear model). The difference between the mean shifts for both groups from both noise conditions was $1.463 \pm 0.511$ kHz ($t(177) = 2.86$ and $p = 0.0051$, $n = 545$ bins, t-test through mixed linear model). Single group data is depicted in Supplementary Fig. 1B, C. One of the groups (Group 1) was further exposed to background noise with an even higher slope (noise #3, green curve in Supplementary Fig. 1A), but did not exhibit a further shift upwards (see Supplementary Fig. 1B). Hence, we focused on noise #2 for all subsequent testing.

**Persistence of spectral centroid shift.** To further assess the robustness of the observed effect, we repeated the experiment in two additional groups of bats (Groups 3 and 4). These two groups were recorded inside acoustic sound-chambers (40–70 dB attenuation) to better control for environmental conditions (Fig. 1b). During the week of silence after noise presentation, we recorded 6314 calls for Group 3 and 3557 calls for Group 4. The calls were summarized and compared to all summarized calls recorded during baseline. Consistent with the results obtained from the first two groups of bats, we found again that the mean centroid of the calls shifted significantly upward relative to baseline ($z = 7.389$ and $p < 0.001$, $5.724 \pm 0.775$ kHz, $n = 292$ bins for Group 3, $z = 3.484$ and $p < 0.001$, $6.65 \pm 1.909$ kHz, $n = 197$ bins for Group 4, z-test through AR(1) model).

In order to compare results across all groups, we looked at either 1) the same time window during which vocalizations occurred (same time of practice) or 2) the same number of vocalizations (same amount of practice) per group. Since calls for Groups 1 and 2 were recorded for a maximum timespan of 1 week after noise cessation, we looked at summarized calls collected during that 1-week period for all groups and compared these to summarized calls recorded during the last week of baseline (1-week data set, Fig. 2a). Similarly, the lowest number of calls recorded for any of the groups was ~5700 calls (Group 1, calls recorded during 1 week after noise cessation). We thus looked at 5000 calls for each condition and for each group (last 5000 baseline calls and the first 5000 calls after the noise perturbation, 5000-data set, Fig. 2b) and generated summarized data sets. All tested groups called consistently during the period after noise cessation (Supplementary Fig. 3). We found that the mean centroid of the calls across all groups, extracted for the summarized 1-week data set, as well as, for the summarized 5000-data set was significantly shifted relative to the bats’ baseline by $3.56 \pm 0.36$ kHz ($t(175) = 9.28$, and $p < 0.001$, $n = 902$ bins, t-test through mixed linear model) and $2.997 \pm 0.468$ kHz ($t(108) = 6.4$, and $p < 0.001$, $n = 568$ bins, t-test through mixed linear model), respectively, and plotted in Fig. 2a, b. Our results suggest that the observed effect was consistent for both time elapsed and amount of practice.

To analyze the stability of the observed centroid shift, we looked at the first and second half of the summarized data sets depicted in Fig. 2a, b. Calls were therefore divided based on their occurrence in the first or second half of the week following the acoustic perturbation or in the first or second half of the first 5000 calls recorded after the acoustic perturbation. This division allowed us to assess the potential decay of the effect as a function of time and amount of practice with the explicit prediction that if the effect is transient, then we would expect the shift to be smaller in the second half when compared to the first half. Interestingly, this was not the case. The mean centroid for the first and second half of the summarized 1-week data set shifted significantly relative to baseline by $2.983 \pm 0.436$ kHz and $4.689 \pm 0.4102$ kHz, respectively ($t(177) = 6.84$ and $p < 0.001$, $n = 882$ bins for the first half, $t(186) = 11.43$ and $p < 0.001$, $n = 944$ bins for the second half, t-test through mixed linear model) and is plotted in Fig. 2c. The mean centroid for the first and second half of the summarized 5000-data set shifted significantly relative to baseline by $2.625 \pm 0.531$ kHz and $3.358 \pm 0.547$ kHz, respectively ($t(111) = 4.94$ and $p < 0.001$, $n = 600$ bins for the first half, $t(105) = 6.14$ and $p < 0.001$, $n = 600$ bins for the second half, t-test through mixed linear model) and is plotted in Fig. 2d. All first and second half data sets were different from each other as well, but were only significant for the half week data sets ($t(131) = 3.77$ and $p < 0.001$, $n = 654$ bins for the half 1-week data sets, $t(82.9) = 1.3$ and $p = 0.198$, $n = 400$ bins for the two half 5000-data sets, t-test through mixed linear model) and did not show a decay but, in fact, a further increase (the difference between the first and second half 1-week data set was $1.712 \pm 0.454$ and $0.738 \pm 0.568$ kHz between the first and second half 5000-data set).

To account for the natural variation of the groups’ social calls, we extracted mean centroid values on a day-by-day basis for each group and compared (across all groups) the difference in the daily value to all other days during baseline and post-noise perturbation periods (Fig. 2e). We found that the values were consistently different for all days between baseline and post-noise perturbation period. Our results thus far suggest that the low-amplitude acoustic perturbation induced a robust form of vocal plasticity across all group compositions and recording environments, which persisted for at least one week.
We further assessed whether the observed changes could be attributed to the fact that these signals were measured during entirely uninhibited, natural interactions which are inherently variable and complex. To do so, we compared the fluctuations of the centroid over a 1-week period during baseline and after-noise exposure for each group (Supplementary Fig. 4). This comparison revealed a clear and persistent separation between the baseline and after-noise curves which never overlapped for any group nor at any time point thus verifying that the observed shift in centroid is not due to random noise in the calls but is an effect of the noise perturbation. In addition, we continued recording Groups 3 and 4 for an extended period of time (5–11 weeks) after the presentation of the noise perturbation to further assess the extended, long-term persistence of vocal plasticity. In detail, Group 3 was recorded for 11 weeks (during which 108970 calls were collected) and Group 4 was recorded for 5 weeks (due to the much lower daily call rate of this group, 16157 calls). Figure 3 shows the change from baseline for each of the two groups, as computed for the mean of the centroid in bins of 5000 calls with an overlap of 500 calls between bins. As expected, and due to the fact that bat social calls are exclusively tied to complex social interactions, the difference from baseline fluctuated somewhat during this extended period of time, yet remarkably, never returned to baseline. Thus, the shift persisted for the entire duration of monitoring, i.e., on the time scale of months.

Analysis of the centroid during the last week of baseline and the week after noise presentation in terms of their variability offers insight into the magnitude of fluctuations. We thus assessed the variability for each group before and after the noise exposure, which revealed that although for two of the four groups the calls differed in their variance in the period after noise perturbation as compared to baseline, the differences for each group were small and not unique in one direction (more or less variable) (Supplementary Fig. 5 and Supplementary Tables 7 and 8). These findings strongly suggest that bats are capable of long-term and persistent vocal plasticity that is not altered in the absence of further changes in the acoustic environment or social constraints.

**Adult vocal plasticity persists across acoustic parameters.** While our manipulation was designed to target changes in the spectral centroid of the bats’ vocalizations, it had no explicit constraints that would prevent the bats from also modifying other features of their calls. To assess this possibility and importantly, to examine the persistence of such potential changes, we computed 10 additional features, which have been used previously to characterize Egyptian fruit bat social calls31. Figure 4a shows the sorted mean shift of these acoustic features for summarized calls measured after the cessation of the acoustic perturbation (comparison was done with summarized calls during the last week of baseline). Consistent with the results presented above, the centroid showed the largest normalized shift. Importantly however, we found that many of the other acoustic features of the bats’ vocalizations also appeared to have changed (Fig. 4a, see Methods, Supplementary Fig. 6 and Supplementary Tables 1–4). To examine whether such a global change might allow us to distinguish the vocalizations of the bats before and after the manipulation, we used an unbiased classifier (Matlab 2017b Classification Learner Toolbox, ensemble classifier Bagged Trees) on the summarized data sets. Indeed, we found that when considering all 11 acoustic parameters (Supplementary Table 4), 86% and 82% of all calls recorded for all groups in the 1-week data sets (419 baseline bins and 483 bins after manipulation), could be correctly assigned as belonging to the 1-week period before and after the noise manipulation, respectively (Fig. 4b). Similarly,
when applying our analysis to the 568 bins (4 groups × 71 bins × 2 conditions) recorded before and after the same manipulation, we found that the unbiased classifier could correctly assign the calls as belonging to the calls recorded before and after the manipulation with accuracies of 83% and 79%, respectively (Fig. 4c).

These results suggest that the groups of bats significantly changed the global acoustic structures of their calls but do not address whether these global changes were also persistent. Figure 4d–g, shows the evolution of each of the 11 acoustic parameters for each group separately during the baseline period (left half of each panel to the dotted red line) and following the cessation of the acoustic perturbation (right half of each panel to dotted red line). For all groups, the divergence of the acoustic parameters after the acoustic perturbation compared to baseline is persistent.

Lastly, and to again account for the natural variation on the groups’ social calls, we extracted normalized mean values on a day-by-day basis for each group and for each of the 11 acoustic parameters and compared (across all groups) the daily means between baseline and the post-noise exposure period (Supplementary Fig. 7). In agreement with the results obtained for the centroid measure (Fig. 2e), we found that for many acoustic parameters the daily post-perturbation period values were significantly different from nearly all baseline days. Together, our results suggest that the changes following the noise perturbation affected the global acoustic structure of bat social calls and importantly, that these global changes were stable and persistent.

Analysis of context-specific vocalizations. Egyptian fruit bat social calls are used exclusively for social interactions and their acoustic content is closely tied to the behavioral context in which they are emitted17. Thus, a potential confound in our results is that the bats did not actually modify the acoustic features of their vocalizations but instead avoided social behaviors that were associated with perturbed vocalizations. To assess this possibility, 13886 calls were recorded in conjunction with video monitoring (a call example and the associated video are depicted in Fig. 5a–c). These calls were manually annotated with respect to the observed behavior associated with each call (Fig. 5c). Annotation was made for calls recorded during the 2 weeks of baseline or for calls recorded during the week of silence after a noise cessation. This type of manual classification was conducted by unbiased observers who were blind to the details and goals of the experiment.

For Group 1, 7868 calls produced after noise #2 and noise #3 were behaviorally annotated and for Group 2, 8018 calls produced during baseline and after noise #2 were behaviorally annotated. Based on previous studies17, the calls were categorized as belonging to six broad behavioral contexts (‘Perch Aggression’, ‘Sleep Aggression’, ‘Food Aggression’, ‘Mating Aggression’, ‘Positive Interaction’, ‘Positive Mating Interaction’). If a behavior could not be identified, it was categorized as ‘Not Classifiable’. Examples of different call types are shown in Supplementary Fig. 2. Calls categorized as ‘Perch Aggression’ and ‘Not Classifiable’ were sorted, summarized and the mean centroid was compared (Fig. 5d, f). For Group 1, the mean centroid of ‘Perch Aggression’ and ‘Not Classifiable’ calls from the period after noise #2 showed the same pattern as previously observed and were shifted relative to baseline by 2.798 ± 0.461 and 1.994 ± 0.591 kHz, respectively. A similar shift was observed for the period after noise #3 where the behaviorally annotated calls were shifted upward by 2.678 ± 0.593 and 2.525 ± 1.03 kHz. When comparing ‘Perch Aggression’ and ‘Not Classifiable’ calls between the two noise conditions, results were not significant relative to each other (z = 0.16 and p = 0.873, n = 136 bins for ‘Perch Aggression’, z-test through AR(1) model; z = 0.298, R = 0.08 and ρ = 0.674, n = 37 bins and 20000 iterations for ‘Not Classifiable’, SMA method, see Methods). When comparing the shift to baseline for the automatic call analysis, we found that the shift for these two noise conditions was 3.118 ± 0.545 and 2.883 ± 0.581 kHz (Fig. 5e) and these were also not significant relative to each other (z = 0.351 and p = 0.725, n = 325 bins, z-test through AR(1) model; Supplementary Fig. 1B). For Group 2, the mean centroids of ‘Perch Aggression’ and ‘Not Classifiable’ calls recorded during the baseline period were elevated by 0.636 ± 0.873 and 1.338 ± 0.781 kHz relative to the noise condition. This slight elevation was caused by calculating the mean over these selected calls which results in a slightly higher centroid in comparison to the mean centroid over all baseline calls from the automatic call analysis. More importantly, these specific types of calls were shifted even higher following the acoustic perturbation by 4.402 ± 0.878 and 3.263 ± 1.35 kHz relative to baseline (Fig. 5f) and the observed shift was significant for the ‘Perch Aggression’ calls (z = 3.388 and p = 0.002, n = 69 bins for ‘Perch Aggression’, z = 1.214 and p = 0.225, n = 138 bins for ‘Not Classifiable’, z-test.

Fig. 3 Stability of centroid shift over weeks and months. Groups 3 and 4 were recorded for an extended period of time after noise cessation (11 and 5 weeks, respectively) to assess the stability of the observed shift in centroid. Depicted is the development of the mean centroid in terms of difference from baseline. A bin number is generated by calculating the mean over 5000 calls and a sliding window of 500 calls. The low call rate for Group 4 resulted in the low bin number for this group. The mean centroid fluctuates for both groups but never returns to zero (− baseline).
In the current study, we investigated the existence of persistent vocal plasticity in adult Egyptian fruit bats (Rousettus aegyptiacus) in response to noise exposure. Without exception, we found that all groups of bats, independent of group composition, environmental conditions and gender, adapted their vocalizations to minimize interference with the acoustic perturbation. Importantly, the observed shift persisted long after noise cessation (in some cases lasting for several weeks or months), which suggests a high degree of vocal plasticity that extends well into adulthood in this species. Our results complement previous studies of vocal plasticity in Egyptian fruit bats that have thus far been restricted to juveniles during development. The demonstration of long-term and persistent vocal plasticity in the adult bat presented in this study further helps establish the Egyptian fruit bat as a feasible and tractable mammalian model organism for studying the neural computations that support vocal plasticity in the adult mammalian brain.

Why do bats retain these unique vocal plasticity abilities as adults? Although this question remains unanswered, it is clear that bats are highly specialized in acoustic signaling. Bats are capable of remarkable vocal control, which up to now has been through AR(1) model). Though the ‘Not Classifiable’ baseline calls are not significantly different to the ‘Not Classifiable’ noise #2 calls, the difference in means follows the same trend as seen for the ‘Perch Aggression’ calls. Furthermore, this effect was comparable to the significant upward shift observed for this group when considering all calls (Fig. 5g, 2.795 ± 0.515 kHz). Combined, analysis of the manually classified calls based on social context reveals a similar pattern of spectral centroid shift, indicating that the observed shifts cannot be accounted for by changes in the social behavior of the group and associated call types. Rather, it is due to a global and stable adaptation of the produced vocalizations.

**Discussion**

In the current study, we investigated the existence of persistent vocal plasticity in adult Egyptian fruit bats (Rousettus aegyptiacus) in response to noise exposure. Without exception, we found that all groups of bats, independent of group composition, environmental conditions and gender, adapted their vocalizations to minimize interference with the acoustic perturbation. Importantly, the observed shift persisted long after noise cessation (in some cases lasting for several weeks or months), which suggests a high degree of vocal plasticity that extends well into adulthood in this species. Our results complement previous studies of vocal plasticity in Egyptian fruit bats that have thus far been restricted to juveniles during development. The demonstration of long-term and persistent vocal plasticity in the adult bat presented in this study further helps establish the Egyptian fruit bat as a feasible and tractable mammalian model organism for studying the neural computations that support vocal plasticity in the adult mammalian brain.
mostly studied in the domain of their “sixth-sense”, i.e., echolocation—a sensory modality used for orientation, navigation and prey acquisition\textsuperscript{35,36}. Although bat species can differ greatly in their lifestyle, foraging strategy and prey items\textsuperscript{37,38}, most of them rely on this extraordinary sensory modality which is highly adaptive in response to changes in their environment. For example, many bats echolocate more loudly, with longer calls and more often or may rapidly shift frequency components of their echolocation calls during increased ambient noise or to avoid jamming by conspecifics\textsuperscript{32,39–44}. These changes, however, are typically short-term, i.e., restricted to the period of noise interference. In contrast, we have found that adult bats adapt the structure of their social calls in response to an external sound source by shifting spectral parameters of their vocalizations and importantly, maintain this shift after noise cessation. The persistence of the shift suggests that vocal plasticity exists beyond short-term changes or adolescence in bats and demonstrates that auditory inputs can drive these changes even in adulthood. The latter notion is consistent with emerging evidence indicating that bat social calls are learned\textsuperscript{5,9,18,19} and might therefore be associated with a high-degree of vocal plasticity that is retained throughout the bat’s life. This notion is ethologically conceivable, as animals who are acoustic specialists might benefit from a more persistent form of vocal adaptation, especially when exposed to an acoustic perturbation affecting the long-term effectiveness of their acoustic communication system.

A major question that remains is what mechanisms are used by bats to modify their vocalizations? According to the source-filter theory of sound production, vocalizations are the products of the vocal tract (tongue, lips, soft palate), which filter source signals originating in the syrinx (in birds) or larynx (in mammals)\textsuperscript{42–47}. As the sound source determines the fundamental frequency through the vibration of the vocal folds, changes in the fundamental are possible via adjusting the sound source while changes in other spectral features occur through adapting the filter of the vocal tract. Comparing all extracted acoustic parameters between baseline and the noise condition in our study revealed that the centroid showed the largest shift, but changes were evident for almost all acoustic parameters. This, in turn, indicates a multifactorial, long-lasting effect induced by the presented acoustic perturbation on the call structure. An animal’s acoustic environment can deviate strongly in its signal-to-noise ratio in a dynamic fashion, enforcing evolved mechanisms that allow the system to deal with these types of variations under noisy conditions\textsuperscript{13,32,43,44,48–51}. Thus, adult bats might retain a high level of control over both the source and filter and can select which aspects of their modification need to be adjusted based on the specific environmental conditions—a notion that is compatible with the stability observed across all extracted parameters.

In addition, it has been postulated that changes in the spectral features of vocalizations could be a by-product of the Lombard effect\textsuperscript{27}—an increase in signal amplitude influenced by noise—where the increase in call amplitude results in the widening of the glottal width and a shortening of the vocal folds shifting spectral features upward\textsuperscript{45}. This, however, is not the cause for the increased centroid in our study since we see no increase in amplitude after noise cessation. On the contrary, the amplitude (calculated from the root mean square) was reduced in comparison to baseline by 1 dB (Supplementary Table 4) and was negatively correlated with the centroid shift, but this correlation is not significant (Supplementary Fig. 9A). The additional measurement of the peak-to-peak amplitude (Supplementary Table 9) showed a similar downward shift after noise perturbation—consistent with the trend found for the RMS—indicating that the call
amplitude did not increase in its peak or average amplitude. A prior study investigating the Lombard effect on echolocation calls of bats found that frequency shifts in response to noise perturbation could occur independently of changes in amplitude44. Nevertheless, the classical definition of the Lombard effect refers to changes in vocalization structure not only in response to an increase in ambient noise levels, but more importantly, during the noise exposure itself48,52. The observed shifts in centroid found in the current study were always measured in complete silence and for an extended period of time after noise cessation. Studies in songbirds36,37 and cetaceans34,35 showing long-term changes in vocalizations in response to noise (over periods of years to decades) have also been linked to the Lombard effect. But, since these studies were conducted in the free field, they did not control for the persistence or disruption of the noise conditions. Consequently, the shifts measured in the current study in silence and for an extended period of time (multiple weeks) after noise cessation cannot be attributed to the Lombard effect, at least not according to its classical definition in the literature. However, it is important to consider that since the Lombard effect is one of many mechanisms underlying vocal plasticity, the underlying mechanisms may not be mutually exclusive.

It has also been shown that habitat or social context can shape vocal signals36–38 (e.g. developing R. aegyptiacus’ social calls can be influenced by calls of an entire social group, known as ‘crowd vocal learning’). Such vocalizations could be termed vocal ‘dialects’19,59 or act as vocal group signatures allowing for group membership and possibly origin to be discerned18,59–63. The bats in the present study all experienced the same noise perturbation and were not exposed to calls by other conspecifics during the recording time, potentially resulting in a group call signature. Thus, when three of the groups had been recaptured after release into a large colony (comprising hundreds of bats), the differences in the measured centroid (consistent with the notion that social calls are highly influenced by acoustic environment) were revealed (Supplementary Fig. 10). It is important to note that the different groups were released to colonies of dramatically different structures in terms of both numbers and sex (e.g. an all-male vs. mixed-gender colony) and future studies in which the colony social structure is characterized and controlled will be important for studying the extent of vocal plasticity in a naturalistic, large-scale social group context.

Together, our results demonstrate persistent vocal plasticity in adult Egyptian fruit bats, thus supporting their potentially important role as a model system for vocal production and learning. Alongside recent technological developments for interrogating the neural circuits of these remarkable mammals, these findings could enable the long-awaited study of the neural circuits supporting vocal plasticity in the mammalian brain.

Methods
Animals. Rousettus aegyptiacus is a megachiropteran, Old-World fruit bat found throughout Africa and the Middle-East. Twenty adult bats were removed from breeding colonies at the University of California, Berkeley, and separated into four groups of six or four bats per group. Group 1 consisted of three males and three females (which gave birth to one pup each toward the end of the experiment), Group 2 of six males, and Groups 3 and 4 of four males each. Each group was housed continuously in a cage with a daily free fly time of 1 h. Food (a fruit-vegetable mix consisting of apples, pears, melons, grapes, plums, kale, carrots and sweet potatoes) and water were given ad libitum. All experimental procedures complied with all relevant ethical regulations for animal testing and research and were approved by the Institutional Animal Care and Use Committee of the University of California, Berkeley.

Group setup. Based on published data by Prat et al.8,17,21 and personal observations we know that these bats do not call in isolation but in groups of bats (>3 animals) where most vocal interactions occur pairwise. Group 1 was chosen to consist of six animals and balanced in gender. During the course of the experiment, all three females became pregnant and gave birth. As the pups’ isolation calls are fairly high in frequency content6,17 the observed upward shift in centroid could have been produced by such calls. Group 2 was then micro-chipped to consist of only males to avoid this potential confound. To test different environmental conditions, Groups 3 and 4 were set in an acoustically quieter and shielded environment (sound boxes, see below). The smaller size of the cages and enclosures used in these experiments limited the number of animals in each group to four (compared to six in the first two groups). Initially, Group 3 and 4 each consisted of three females each, but as both female groups barely communicated vocally (<80 calls triggers per day) already during the baseline period, these groups were discontinued before any noise manipulation and instead rearranged to consist of only males (four individuals per group). Importantly, none of the bats had experienced any type of playback prior to this study.

Experimental setup. Groups 1 and 2 were housed in a metal cage (81 × 48 × 94 cm) with a wire spacing of 1.3 cm set at an overall height of 163 cm (Fig. 1A) and positioned in a corner of a larger housing room. The cage was surrounded by felt curtains to dampen echoes. Two loudspeakers (Discovery Tweetter R2604/833000, Scanspeak, Videbæk, Denmark) were positioned at a height of 120 cm and 10 cm distance with respect to the front and the back of the cage. At the front of the cage, the microphone (custom built from Knowles SPU4101-RSH-QR, Itasca, IL, USA) was placed at a distance of 10 cm and height of 120 cm. An additional USB camera (Megasipix IP camera, ELP, Allipu Technology Co. Ltd, Shenzhen, Guangdong, China) was mounted to a side of the cage later in the course of the experiment and used to record (motion-based, triggered video recordings) the events in the cage. Group 2 was housed in a plastic mesh cage (51 × 55 × 30 cm) in an acoustic chamber (44 × 76 × 60 cm) lined with acoustic foam, (Fig. 1B). The microphone was positioned in a back corner 10 cm from the ceiling, pointing towards the cage, and the speaker was mounted in the middle of the ceiling facing backwards towards the cage. A USB camera was then placed in the middle of the cage door with the cage being in its field of view. Microphones were connected directly and the speakers via a stereo amplifier (Servo 120 A, Sanomax Technologies, Hicksville, NY, USA) to an A/D-D/A converter (UltraLite-mk3, MOTU, Cambridge MA, USA) with a sampling rate of 192 kHz, which sent and received its signals from a PC. Recordings were triggered through a combination of signal detection and loudness. Recording and playback was controlled via the Soundmexpro (HorTech gGmbH, Oldenburg, Germany) toolbox and Matlab (Matlab R2011a, MathWorks, Natick, MA).

Procedure. Each group of bats was monitored for 1 week while they became accustomed to the new environment and social arrangement. After this habituation, their vocal activity was recorded for 2 weeks to establish a baseline call behavior. Following this time period, playback of continuous background noise (i.e. noise perturbation) commenced. The bats were exposed for up to 2 weeks to a background noise followed by 1 week of silence. This was repeated for up to two or three different noise conditions for Groups 2 or 1, respectively.

Stimuli. Different noise conditions were defined through their spectral tilts (Fig. 1C and Supplementary Fig. 1A). This was accomplished by designing an impulse response with a defined level decrease per octave (−3.−6, and −12 dB per octave); a steeper downward slope results in less activation of higher frequency auditory fibers. The stimulated level resulted in three different noise conditions (noise #1), with a shallower slope and the most energy between 8–27 kHz, the second noise condition (noise #2), with a steeper slope and its energy shifted towards lower frequencies (4–20 kHz) and the third noise condition (noise #3), with the steepest slope and energy mainly in the low frequency range (below 5 kHz). By convolving white noise with the impulse response, the spectral tilt is imposed on the presented noise. The speaker’s frequency response was not corrected as it matched the noise’s spectral shape. The background noise was presented at 45–50 dB SPL (as used by Luo and Wiegrebe25) and was continuously active for the two playback weeks. Groups 1 and 2 first experienced noise #1 followed by noise #2. Group 1 was furthermore exposed to noise #3. Groups 3 and 4 were only exposed to noise #2. For Groups 1 and 2, noise was silenced for 1 week between the two different noise presentations (noise #1 followed by noise #2) to observe long-term effects on the bats’ vocalizations. After the last noise condition and the subsequent week of silence, Groups 1 and 2 were released back to their original colony. Before releasing them to the colony, the groups were exposed for an additional 11 or 5 weeks, respectively, to measure whether the bats would adapt their calls back to their original baseline or retain their adapted calls. Group 4 was released after 5 weeks due to a low call rate and unchanged call parameters. Group 3 was released after 11 weeks with no apparent reversal in call parameters to their baseline. Groups 2, 3, and 4 were recaptured after being back in the colony for 1 month, brought back to their previous experimental setups and recorded for 1 month. The purpose was to investigate whether the bats would adapt their calls back to their original baseline after being exposed to a more natural acoustic environment.

All noise conditions were played via the setup speaker(s) and recorded with a calibrated condenser microphone (¼” piezoelectric free-field microphone, ¼” CCP preamplifier 26CB, G.R.A.S. Sound & Vibration A/S, Holte, Denmark) with a flat frequency response up to 100 kHz to ensure correct frequency presentation of the
noise. Amplitude levels were adapted with the help of a sound calibrator (1 kHz at 94 dB, Tenma 72–7260, Leeds, UK).

**Call detection and analysis.** A Matlab-based, custom designed code automatically cut sound files into either cage-related noise in the recordings. Recordings were low-pass filtered and peaks in the sound envelope above a certain threshold were detected, resulting in an initial signal cut. Duration and root mean square thresholds determined whether a detected signal was a call or not. An additional custom designed function evaluated the envelope of the detected signal which was checked against a duration threshold for a second time. Consequently, detected calls were filtered (with frequency limits set to 0.5 and 10 kHz) and then analyzed according to their duration, root mean square (RMS), entropy, peak frequency, minimum and maximum frequencies, frequency band width, centroid, fundamental, Wiener and Shannon entropy. The peak frequency was detected in the call’s magnitude spectrum, the minimum and maximum frequencies were at ±2.5 dB cutoff points of the peak frequency, and the frequency bandwidth was the difference between these two cutoff points. The centroid, fundamental and Wiener and Shannon entropy were calculated on the median over all call snippets (10% of entire call length) and a sliding window of 9% of the entire call length over the entire call. The centroid was measured by dividing the sum of a set frequency range (for Groups 1 and 2, 0.5–0.6 kHz and Groups 3 and 4, 0.5–0.5 kHz) by the sum of the power spectrum of the frequency range. For extraction of the fundamental the Matlab-based functions ‘spCorr’ and ‘spPitch-Corr’ were adapted (with frequency limits set to 0.5 and 10 kHz). For evaluation of the Wiener entropy, the Matlab-based function ‘spectral flatness’ was used and for the Shannon entropy and the entropy (calculated over the spectrum and the time signal) the Matlab-based function ‘entropy’ was employed. The RMS was the root mean square of the call amplitude in voltage. For further analysis, only calls with a centroid below 80 kHz and a fundamental between 0.5 and 5 kHz were considered valid, as this reduced the risk of including cage noise.

**Data summarization.** As the Lilliefors test revealed that the data sets were not Gaussian distributed at a 5% significance level, we applied a summarization method on data sets. This was done by averaging every 70 calls, resulting in a number of observation bins (e.g. 6348 baseline calls for Group 1 resulted in 90 observation bins, each consisting of a mean over 70 calls). The resulting errors of these means are closer to a normal distribution than when calculated over the non-summarized data. The Lilliefors test on the summarized data verified its normal distribution. This summarization into observation bins was applied to each condition of the experimental data (baseline and after noise manipulation) and group. Due to overall smaller sample sizes of the half data-sets (of the 1-week and 5000 calls) and the manually classified calls we averaged over 50 calls and 35 calls, respectively, to ensure at least about 30 observation bins for each data set (except for one exception see below). Statistics (see below) was applied on summarized data sets.

For normalization, each group data was normalized by first subtracting from the mean of the baseline observation bins and then dividing by the maximum of the absolute value for the baseline observation bins.

**Statistics.** When comparing baseline and post-perturbation conditions within one group, we applied a z-test for comparing between two time-series; this test is described in section 15.2 of Ramsey and Schafer. The test assumes each time series has a constant mean, that observations within the time series exhibit 1st order autocorrelation (AR(1)), and that the autocorrelation and variance is the same for both time series. We refer to this test as the ‘z-test through AR(1) model’.

For each group we reported the estimated AR(1) correlation, $r_A$, and pooled standard deviation in Supplementary Tables 5 and 6, and the differences of the least square means and corrected standard errors (mean ± corrected standard error) in the main results section.

The smaller sample size of the video annotated data set for Group 1 ‘Not Classifiable’ calls resulted in less than 30 observation bins per noise #2 and noise #3 condition. To statistically analyze these short, autocorrelated data sets we applied the Simulation Modelling Analysis (SMA) developed by Boscardin and Nash to calculate $r_A$ for small sets of time-series data collected over time. In brief, SMA first estimates the means of each condition and a common AR(1) correlation, $r_A$, then, under a null-model of no difference in means, simulates many datasets under the original data structure, with observations having $r_A$ correlation. The resulting empirical p-value is the proportion of simulated datasets having differences in means that were at least as extreme as the originally observed difference in means.

For comparing means between baseline and post-perturbation conditions from multiple groups, we applied a mixed linear model, which assumed observations within each group-condition combination had a common variance and AR(1) correlation. We estimated the denominator degrees of freedom for the resulting t-tests according to the Kenward-Roger method. For each time-period and group we report the degrees of the least square means and the standard errors in the main results section.

For each parameter a linear regression extracted the slope of the measure over time for baseline and noise #2. The differences between baseline and noise #2 slope for each parameter was fitted by a randomized complete block ANOVA test where group by time-period parameter a fixed factor. This analysis allowed us to test the natural relationship between baseline and noise #2 slopes and for the correlation of parameters from the same group.

All values were obtained by combining results from the SAS statistical software (SAS University edition, SAS Institute Inc., Cary, U.S.A.) or from the SMA free commercial software (Version 11.10.18, Copyright J. B. Boscardin, with customized Matlab codes. Additionally, the alpha level of 5% is Bonferroni corrected to account for multiple comparisons (11 call parameters → 0.05/11 = 0.0045).

**Classifier.** The Bagged Trees classifier in the Matlab classification Learner toolbox uses the Breiman’s ‘random forest’ algorithm (for reference, see Breiman et al.). However, the classifier assumes independence among all observations and the observed 1st order autocorrelation for each parameter data set is between 0.206 and 0.523.

**Behavioral analysis.** Group 1 calls recorded during the week of silence after exposure to noise #2 and #3 and Group 2 calls recorded during the baseline period and during the week of silence after noise #2, were selected for annotation based on behavior observed in the videos by student volunteers. These unbiased observers were blind to the specifics of the project design and goal. A valid annotation meant that a call could be reliably synchronized with a video sequence and the behavioral content identified. Due to the camera’s field of view, a select few identifiable behaviors were chosen for annotation. These behavior types were based on a previous behavioral classification study done for the same species (for details see Prat et al.).

**Perch aggression:** aversive behavior of one bat to another when in close vicinity to each other.

**Sleep aggression:** aversive behavior when at least one bat had been sleeping.

**Food aggression:** squabbles between bats with food involved.

**Mating aggression:** aversive behavior between a male and female during mating.

**Positive interaction:** non-aversive behavior between bats, typically during grooming.

**Positive mating interaction:** non-aversive behavior between bats during mating.

The goal of this manual analysis was to verify that any change in the bats’ vocalization parameters was not due to a change of call type induced by a change in behavior. If the call type could not be determined it was termed ‘Not Classifiable’. Due to nature of the setup and recording procedure, a large amount of the audio recordings was cage-related noise. The manual analysis of the calls therefore additionally provided proof that only a small percentage of cage-related noise contaminated the automatic post call analysis.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

All data supporting the findings in this study is available from the corresponding author upon request.

**Code availability**

All code designed for analyzing the data in this study is available from the corresponding author upon request.

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**References**

1. Guenther, F. H. A neural network model of speech acquisition and motor equivalent speech production. *Biol. Cybern.* 72, 43–53 (1994).

2. Callan, D. E., Kent, R. D., Guenther, F. H. & Vorperian, H. K. An auditory-feedback-based neural network model of speech production that is robust to developmental changes in the size and shape of the articulatory system. *J. Speech, Lang. Hear. Res.* 43, 721–736 (2000).

3. Oller, D. K. & Eilers, R. E. The role of audition in infant babbling. *Child Dev.* 59, 441–449 (1988).

4. Reichmuth, C. & Casey, C. Vocal learning in seals, sea lions, and walruses. *Carr. Opin. Neurobiol.* 26, 66–71 (2014).

5. Belov, C. I. & Jarvis, E. D. Birds, primates, and spoken language origins: behavioral phenotypes and neurobiological substrates. *Front. Evol. Neurosci.* 4, 12 (2012).

6. Dreu, A. K., Tieleman, D. P., Kuhl, P. K. Birdsong and human speech: common themes and mechanisms. *Annu. Rev. Neurosci.* 22, 567–631 (1999).

7. Stoeger, A. S. & Manger, P. Vocal learning in elephants: neural bases and adaptive context. *Carr. Opin. Neurobiol.* 28, 101–107 (2014).

8. Prat, Y., Taub, M. & Yovel, Y. Vocal learning in a social mammal: Demonstrated by isolation and playback experiments in bats. *Sci. Adv.* 1, e1500019 (2015).

9. Doupe, A. J. & Kuhl, P. K. Birdsong and human speech: common themes and mechanisms. *Annu. Rev. Neurosci.* 22, 567–631 (1999).

10. Reichmuth, C. & Casey, C. Vocal learning in seals, sea lions, and walruses. *Carr. Opin. Neurobiol.* 26, 66–71 (2014).

11. Prat, Y., Taub, M. & Yovel, Y. Vocal learning in a social mammal: Demonstrated by isolation and playback experiments in bats. *Sci. Adv.* 1, e1500019 (2015).
19. Prat, Y., Azoulay, L., Dor, R. & Yovel, Y. Crowd vocal learning induces vocal production learning in bats. *Anat. Rec. - Part A Discov. Mol., Cell., Evolut. Biol.* 287, 1080–1102 (2005).

20. Verner, S. C. What bats have to say about speech and language. *Science* 354, 145–149 (2016).

21. Neuwiler, G. The Biology of Bats. (Oxford University Press, Oxford United Kingdom 2000).

22. Denzerger, A. & Schnitzius, H. U. Bat guards, a concept to classify the highly diverse foraging and echolocation behaviors of microchiropteran bats. *Front. Physiol.* 4, 164 (2013).

23. Gross, D., Yovel, Y. & Tarsyev, M. M. Neuroethology of bat navigation. *Curr. Biol.* 28, R997–R1004 (2018).

24. Amichai, E., Blumrosen, G. & Yovel, Y. Calling longer and louder: how bats use biosonar under severe acoustic interference from other bats. *Proc. R. Soc. B Biol. Sci.* 282, 20152064 (2015).

25. Yartsev, M. M., Witter, M. P. & Ulanovsky, N. Grid cells without theta oscillations in the entorhinal cortex of bats. *Nature* 479, 103–107 (2011).

26. Yartsev, M. M. & Ulanovsky, N. Representation of three-dimensional space in the Hippocampus of flying bats. *Science* 340, 367–372 (2013).

27. Brumm, H. & Slabbekoorn, H. Acoustic communication in noise. in *Principles of animal communication* (eds E. G. Fitch, S. N. & Fischer, K. E.) Mammalian aging, metabolism, and ecology: evidence from the bats and marsupials. *J. Gerontol.* 46, B47–53 (1991).

28. Kwiecinski, G. G. & Griffiths, T. A. Rousettus egiptiacus. *Mamm. Species* 611, 1–9 (1999).

29. Zhang, W. & Tarsyev, M. M. Correlated Neural Activity across the Brains of Socially Interacting Bats. *Cell* https://doi.org/10.1016/j.cell.2019.05.023 (2019).

30. Prat, Y., Taub, M. & Yovel, Y. Everyday bat vocalizations contain information about emitter, addressee, context, and behavior. *Sci. Rep.* 6, 39419 (2016).

31. Iversen, S. C. What bats have to say about speech and language. *Science* 354, 145–149 (2016).

32. Luo, J., Goerlitz, H. R., Brumm, H. & Wiegrebe, L. Linking the sender to the receiver: vocal adjustments by bats to maintain signal detection in noise. *Science* 5, 18555 (2015).

33. Tessler, J. & Shommerman, M. S. Context-dependent effects of noise on echolocation pulse shape and period. *Annu. Rev. Neurosci.* 36, 489–517 (2013).

34. Turner, E. C. & Brainard, M. S. Performance variability enables adaptive plasticity of ‘crystallized’ adult birdsong. *Nature* 450, 1240–U11 (2007).

35. Reiner, A., Yamamoto, K. & Karten, H. J. Organization and evolution of the avian forebrain. *Anat. Rec. - Part A Discov. Mol., Cell., Evolut. Biol.* 287, 1080–1102 (2005).

36. Neuweiler, G. The Lombard effect and other noise-induced vocal modifications: insight from mammalian communication systems. *Biophys. Rev.* 88, 809–824 (2013).

37. Brumm, H. & Slabbekoorn, H. Acoustic communication in noise. in *Advances in the Study of Behavior*, Vol 35 (eds Slater, P. J. B., Snowdon, C. T., Brockmann, H. J., Roper, T. J. & Naguib, M.) 1–209 (Academic Press, Cambridge, MA 2003).

38. Slabbekoorn, H. Some of the city: noise-dependent spectral plasticity in the acoustic phenotype of urban birds. *Anim. Behav.* 85, 1089–1099 (2013).

39. Kozy, G., Hefner, R. S. & Hefner, H. E. Hearing in a megachiropteran fruit bat (Rousettus egiptiacus). *J. Comp. Physiol. Psychol.* 112, 371–382 (1998).

40. Prat, Y., Taub, M., Pratt, E. & Yovel, Y. Data descriptor: an annotated dataset on bat vocalization in a non-songbird. *Curr. Exp. Biol.* 3, 371–373 (2013).

41. Hage, S. R. & Metzner, W. Potential effects of anthropogenic noise on echolocation behavior in horseshoe bats. *Commun. Integr. Biol.* 6, 24753–24753 (2013).
Author contributions
D.G. and M.M.Y. designed the study, analyzed the data, and wrote the paper. D.G. conducted the experiments, J. D and E.P annotated the video data. All authors discussed the results and commented on the paper.

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