



Original Article

The effects of stress and glucocorticoids on vocalizations: a test in North American red squirrels

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Received 31 October 2018; revised 20 February 2019; editorial decision 5 March 2019; accepted 16 March 2019; Advance Access publication 10 April 2019.

Acoustic signaling is an important means by which animals communicate both stable and labile characteristics. Although it is widely appreciated that vocalizations can convey information on labile state, such as fear and aggression, fewer studies have experimentally examined the acoustic expression of stress state. The transmission of such public information about physiological state could have broad implications, potentially influencing the behavior and life-history traits of neighbors. North American red squirrels (*Tamiasciurus hudsonicus*) produce vocalizations known as rattles that advertise territorial ownership. We examined the influence of changes in physiological stress state on rattle acoustic structure through the application of a stressor (trapping and handling the squirrels) and by provisioning squirrels with exogenous glucocorticoids (GCs). We characterized the acoustic structure of rattles emitted by these squirrels by measuring rattle duration, mean frequency, and entropy. We found evidence that rattles do indeed exhibit a “stress signature.” When squirrels were trapped and handled, they produced rattles that were longer in duration with a higher frequency and increased entropy. However, squirrels that were administered exogenous GCs had similar rattle duration, frequency, and entropy as squirrels that were fed control treatments and unfed squirrels. Our results indicate that short-term stress does affect the acoustic structure of vocalizations, but elevated circulating GC levels do not mediate such changes.

Key words: bioacoustics, red squirrel, stress, vocalization.

INTRODUCTION

Acoustic communication is a critical means by which information is transferred within and among animal species. Vocalizations can convey stable information on various characteristics of signalers, such as individual identity (Beer 1970; Beecher 1989; Blumstein and Munos 2005), body weight and size (Clutton-Brock and Albon 1979; Fitch 1997; Bee et al. 1999; Reby and McComb 2003; Blumstein

and Munos 2005; Koren and Geffen 2009), sex (Blumstein and Munos 2005; Ey et al. 2007), and social rank (Clark 1993; Muller et al. 2004; Koren et al. 2008; Yosida and Okanoya 2009; Terleph et al. 2016), and they are often encoded with several layers of information, for example, rank, sex, and individual identity (Koren and Geffen 2009). Communicating this information is consequential for both signalers and receivers, serving a wide array of functions, from attracting mates (Andersson 1994) to reducing conflict and maintaining affiliations in social groups (Masataka and Symmes 1986; Digweed et al. 2007; Soltis et al. 2005a).

Vocalizations can also contain information on labile traits, such as short-term stress state or the changes in glucocorticoids (GCs) that

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are released in response to an acute environmental challenge. Here, we differentiate short-term stressors as discrete events that stimulate an increase in GCs that lasts just minutes, from chronic stressors, that stimulate a continued release in GCs over the course of days or longer. GCs, a class of steroid hormones secreted by the adrenal glands, are released shortly after a stressful event and perform an array of functions in mediating an organism's physiological stress response, including enhancing the effects of the first wave of response from hormones such as epinephrine and norepinephrine (Sapolsky et al. 2000). Stress is known to influence the acoustic structure of vocalizations in a number of species (Manser 2001; Sacchi et al. 2002; Slocombe et al. 2009; Wilson and Evans 2012). Motivation-structural rules make predictions about the characteristics of vocalizations produced in high-stress contexts: hostile vocalizations tend to be lower in frequency and noisier (highly entropic), and fearful vocalizations tend to be higher in frequency and more tonal (Morton 1977). Although some studies have found empirical support for these rules, others have found inconsistencies. For example, vocalizations associated with fear often fail to consistently conform to these motivation-structural rules and are often highly entropic (Morton 1977; August and Anderson 1987). The effects of short-term stress on vocalization structure are thus difficult to generalize.

Although many studies have examined the structure of vocalizations produced in high-stress situations, they have concentrated primarily on vocalizations produced in just a few contexts, and most of them have been observational (Morton 1977; Biben et al. 1986; Zuberbuhler 2009). Most studies have focused on social contexts, including calls produced by victims in agonistic encounters between social group members (Morton 1977), alarm calls (Zuberbuhler 2009), separation between mothers and their young, and between social group members (Biben et al. 1986; Bayart et al. 1990; Rendall 2003; Ehret 2005). Other research has centered on begging calls (Sacchi et al. 2002; Perez et al. 2016) and distress screams produced by individuals in imminent danger of predation or of being seized by a predator, which likely function to solicit intervention from another animal capable of interfering in social species (Hogstedt 1982; Lingle et al. 2007; Blumstein et al. 2008).

Very few studies have experimentally examined the influence of stress or changes in GCs on vocalization structure. One notable exception is Perez et al. (2012), who assessed the effects of GCs on the acoustic structure of zebra finch (*Taeniopygia guttata*) vocalizations. Their experiment included 2 stress treatments: social isolation and treatment with exogenous GCs, and they found that both types of stress significantly altered vocalization features. Compared with untreated individuals, zebra finches in both treatment groups emitted vocalizations of higher frequency than those in the control group (Perez et al. 2012).

The literature on the influence of stress on vocalizations skews heavily toward group-living species and focuses primarily on just a few contexts in which stress occurs; far less is known about the relationship between stress and vocalization structure in solitary species, despite the fact that many regularly produce vocalizations in short-term stress-inducing situations (Hogstedt 1982). Furthermore, few studies have experimentally examined this relationship, leaving a gap in our understanding of the mechanism by which stress may influence acoustic structure. We examined how a short-term stressor (resulting from trapping and handling) and administration of exogenous GCs affected the territorial vocalizations of solitary, territorial North American red squirrels (*Tamiasciurus hudsonicus*).

Red squirrels defend discrete territories throughout the year and produce vocalizations called “rattles” that advertise territorial

ownership (Smith 1968), which deters intruders (Siracusa et al. 2017). At the center of each territory is a “midden,” a network of underground tunnels that serves as storage space for white spruce (*Picea glauca*) cones that compose 50–80% of a squirrel's annual diet (Donald et al. 2011; Fletcher et al. 2013). Overwinter survivorship without a midden is near zero (Larsen and Boutin 1994). Successful defense of a territory against pilferage from the midden, therefore, represents an important component of overwinter survival for a red squirrel.

Red squirrel rattles contain stable information on individual identity (Digweed et al. 2012; Wilson et al. 2015), and receivers discern encoded kinship information, though this may be context dependent (Wilson et al. 2015; Shonfield et al. 2017). In a playback experiment, focal squirrels only differentiated between the rattles of kin and nonkin when the playback rattles used were emitted by squirrels that had just been live-trapped and handled (henceforth, “post-trap rattles”) (Shonfield et al. 2017). This stress-related context dependency of kin discrimination indicated that possible differences in acoustic structure of stressed and nonstressed squirrels warranted examination.

To test this directly, we conducted a 2-part study to examine the relationship between stress state and rattle acoustic structure. In the first experiment, we recorded rattles of wild red squirrels after they were live-trapped and handled and compared these to rattles recorded opportunistically, without provocation from squirrels moving freely around their territories. Previous studies verified this method of inducing stress: squirrels exhibit a substantial increase in circulating GC levels minutes after entering a trap and during handling (Bosson et al. 2012; van Kesteren et al. 2018 preprint).

To identify if elevated circulating GCs are part of the mechanism by which a short-term stressor (such as capture and handling) alters rattle acoustic structure, we conducted a second experiment where we treated squirrels with GCs (dissolved in a small amount of food) and compared their rattles to those of squirrels in a positive control group (provided with the same amount of food but without GCs) and a negative control group (provided with no food or GCs). A previous study showed that in GC-treated squirrels, plasma GCs rose quickly after treatment and then slowly declined over the ensuing 12 h (van Kesteren et al. 2018 preprint).

We first predicted that if rattles do encode information about stress state and recording settings are consistent across conditions, post-trap rattles would be structurally distinct from opportunistic rattles. Based on the results of Perez et al.'s (2012) zebra finch experiments, we predicted that post-trap rattles would be higher in frequency. We then predicted that if GCs are the mechanism by which short-term stress alters rattle acoustic structure, rattles emitted shortly after treatment with exogenous GCs would exhibit the same structural distinctions as post-trap rattles when compared with rattles produced prior to treatment and rattles produced by positive control and negative control squirrels over the same period of time. We expected these structural distinctions to be graded, peaking shortly after treatment and then declining as a function of time since consumption of treatment mirroring the peak and decline of circulating GC levels following treatment.

METHODS

Study site and species

This study was part of the Kluane Red Squirrel Project, a long-term study of a wild population of red squirrels that has been

tracked continuously since 1987 (McAdam et al. 2007) within Champagne and Aishihik First Nations traditional territory in the southwestern Yukon, Canada (61°N, 138°W). The habitat is an open boreal forest dominated by white spruce trees (*Picea glauca*; Krebs et al. 2001). All squirrels were marked individually with ear tags with distinct alphanumeric combinations (Monel #1; National Band and Tag, Newport, KY), and wires in unique color combinations were threaded through the ear tags to allow for individual identification from a distance. We livetrapped squirrels periodically to track female reproductive state and territorial ownership using tomahawk traps (Tomahawk Live Trap Company, Tomahawk, WI) baited with peanut butter (McAdam et al. 2007).

Trap-induced stress experiment field methods

We collected rattles from squirrels across 4 study areas between April and August in 6 separate years from 2005 and 2017 (Table 1). In the capture-induced stress experiment, we compared the structure of rattles collected opportunistically to rattles collected shortly after a squirrel was trapped, handled, and released ("trap rattles"). We collected rattles for this experiment using a Marantz digital recorder (model PMD 660; 44.1 kHz sampling rate; 16-bit amplitude encoding; WAVE format) and a shotgun audio recorder (Sennheiser, model ME66 with K6 power supply; 40–20 000 Hz frequency response [± 2.5 dB]; super-cardioid polar pattern). To collect opportunistic rattles, we stood on a squirrel's midden at a distance of no greater than 5 m from the squirrel until it produced a rattle. To collect trap rattles, we trapped and handled a squirrel on its midden, then recorded its first rattle on release from a handling bag (within a minute of release). Trapping and handling is part of ongoing research activity; it was not done explicitly for this study. Human presence was controlled for: the same person that trapped and handled the squirrel also recorded the rattle at a distance no greater than 5 m from the squirrel; thus, only 1 person was present for the recording in both treatment conditions. Red squirrels rattle spontaneously and in response to detection of conspecifics (Smith 1978), but we cannot rule out the possibility that the rattles were elicited by the person recording. However, even if squirrels were rattling in response to the presence of a person, because the stimulus was the same in both conditions (only 1 person was present), any differences in call structure could not reflect differences in the stimulus, and thus the simplest explanation is that any differences in call structure reflect the difference in physiological stress state prior to the arrival of the stimulus. We did not record the exact amount

of time a squirrel spent inside of a trap, but squirrels were in traps for no more than 120 min before they were released and a rattle was collected. As would be expected, squirrels exhibit a substantial increase in circulating GC levels minutes after entering a trap and during handling (Bosson et al. 2012; van Kesteren et al. 2018 preprint).

In total, 351 rattles from 235 unique individuals (308 opportunistic rattles from 205 squirrels, 39 post-trap rattles from 30 squirrels) were recorded and analyzed in the years 2005, 2006, 2009, 2015, and 2016. Of the 235 squirrels, 127 were male and 108 were female (Table 1). These rattles were part of a long-term data set of rattles compiled by prior researchers with the Kluane Red Squirrel Project.

Exogenous GC treatment experiment field methods

In the second experiment, we assessed the influence of experimental increases in circulating GCs on rattle acoustic structure. We sought to track graded changes in rattle acoustic structure over an extended period of time induced by the GC treatment instead of a simpler pretreatment/post-treatment analysis. We compared the rattles of squirrels in 3 treatment groups, using an established protocol for oral administration of GCs. In the experimental group ($n = 16$), individuals were fed 8 g of peanut butter (all natural, no sugar, salt, or other additives) mixed with 2 g of wheat germ and 8 mg of cortisol (hydrocortisone, Sigma H004). This treatment causes a significant increase in circulating GCs, peaking 90–120 min post-treatment (Dantzer et al. 2013; van Kesteren et al. 2018 preprint). Individuals in the positive control group ($n = 16$) were fed the same amount of peanut butter and wheat germ with no cortisol added. Each squirrel in these 2 treatment groups was treated for 1 day (see details below). Lastly, we had a negative control group of squirrels that were not fed or manipulated in any way ($n = 23$). Our experimental positive control group and our negative control group (the latter of which live on a nearby study area) were comprised exclusively of male squirrels. However, no sex differences are known to exist in rattle acoustic structure (Wilson et al. 2015).

The morning of treatment, between 0730 and 1000 h, for each squirrel in the GC-treated and positive control groups, we placed one treatment in a bucket hanging in a tree near the center of its midden. Pilferage from buckets was extremely low (van Kesteren et al. 2018 preprint), ensuring that treatments were eaten by the target squirrel and not neighboring conspecifics or heterospecifics.

Table 1
Number of rattles collected by year, study grid, collection method, and date range

Year	Grid: AG	Grid: KL	Grid: SU	Grid: JO	Collection method	Date range
2005	0	2 (2,0)	3 (1,2)	0	Opportunistic: 1 Post-trap: 4	7 June–31 July 2005
2006	0	113 (66,47)	93 (43,50)	0	Opportunistic: 204 Post-trap: 2	13 June–14 July 2006
2009	30 (15,15)	53 (26,27)	8 (6,2)	0	Opportunistic: 54 Post-trap: 37	26 March–26 July 2009
2016	24 (12,12)	25 (14,11)	0	0	Opportunistic: 49 Post-trap: 0	6 June–2 August 2016
2017	0	93 (93,0)	22 (22,0)	599 (599,0)	Zoom mic: 714 - Neg Control: 115 - Pos Control: 367 - GC: 232	2 June–14 August 2017

In parentheses, rattles are split up by sex (male, female). For some squirrels, more than one rattle was collected.

We recorded the time each treatment was placed in each bucket and checked the buckets throughout the morning at a minimum of once every hour and maximum of every 45 min in order to determine the 1-h time frame in which the squirrel consumed its treatment. Peanut butter treatments were delivered to the focal individuals' buckets in paper cups; consumption of treatment was confirmed by finding an empty paper cup in their bucket. As a part of another study (Dantzer et al. 2013), we have been providing squirrels with supplemental peanut butter for >10 years and have never observed squirrels caching peanut butter. All treatments were consumed between 0830 and 1130 h. Eight squirrels (positive control, $n = 4$; GC, $n = 4$) did not consume their treatments by 11:30 h; these treatments were removed from the bucket and the squirrels were excluded from analyses. Two individuals (GCs $n = 2$) consumed their treatment over a period of several hours instead of consuming it within a 1-h time block. Because we sought to simulate short-term stress induced by a rapid elevation of circulating GC levels, these squirrels were excluded from analysis as well. Our final sample size was GC ($n = 10$), positive control ($n = 12$), and negative control ($n = 23$).

We recorded rattles using stationary Zoom H2N Audio Recorders (Zoom Corporation, Tokyo, Japan) that were covered with windscreens and attached to 1.5-m stakes in the center of each squirrel's midden. Because they are not weather proof, we placed an umbrella 30 cm above each audio recorder to protect it from harsh weather conditions. We set the audio recorders in 44.1 kHz/16 bit WAVE format and recorded in 2-channel surround mode. We deployed the audio recorders between 1700 and 2200 h on the day before treatment so that they would collect "pretreatment" rattles the following morning prior to treatment. They recorded continuously until nightfall on the day of treatment, recording rattles of the target squirrel, neighboring individuals, and other ambient noise. Rattles recorded in the evening prior to treatment were excluded from analysis; thus, all rattles analyzed in this experiment were recorded on the day of treatment, between approximately 0600 and 2330 h. We chose this recording period because this recording window should have captured rattles at natural GC levels (pretreatment rattles), during the post-treatment spike in circulating GC levels, and the ensuing decline. This is based on our previous study showing that when squirrels are fed exogenous GCs, plasma cortisol concentrations spike within 90–120 min of treatment and decline over the ensuing 12 h (van Kesteren et al. 2018 preprint).

In order to analyze rattles recorded on stationary zoom recorders, we used Kaleidoscope software (version 4.3.2; Wildlife Acoustics, Inc., Maynard, MA) to detect rattles in the recordings. Detection settings were: frequency range: 2000–13 000 Hz; signal duration: 0.4–15 s; maximum intersyllable silence: 0.5 s; fast Fourier transform size: 512 points (corresponding to a temporal resolution of 6.33 ms and a frequency resolution of 86 Hz); distance setting: 2 (this value ensures that all detections are retained). Previous research using our same population, recording apparatus, and rattle extraction technique, and ground-truthed by comparing the results to those obtained by a human observing the squirrels being recorded, showed that our method detects not only 100% of a focal squirrel's rattles (see Siracusa et al. forthcoming) but also detects nonrattles and the rattles of neighbors.

We used a previously developed technique for distinguishing focal squirrel rattles from nonrattles and neighbor rattles (Siracusa et al. forthcoming). We first automatically analyzed the acoustic structure of every detection using the R package "Seewave" (version 2.0.5; Sueur et al. 2008). Structural features included duration,

root mean square amplitude, pulse rate, duty cycle, peak frequency, first energy quartile, skewness, centroid, and spectral flatness (see detailed definitions in Sueur et al. 2008 and Siracusa et al. forthcoming). We analyzed a more complex suite of rattle characteristics here because these features encode the most information about individual identity (Digweed et al. 2012). Second, we used SPSS (software, version 24, IBM Corporation, Armonk, New York) to apply a previously established linear discriminant function analysis model to the structural measurements of each detection. The model, which was developed during the same ground-truthing experiment described above, labeled each detection as "focal rattle," "neighbor rattle," or "nonrattle," and assigned a probability that the detection was a focal rattle. Third, we used Kaleidoscope to review spectrograms of all detections labeled "focal rattle" that have an estimated probability of being a focal rattle of at least 0.999. During this step, we removed any nonrattles that were included erroneously as focal rattles.

Our final data set included 714 rattles from 45 focal squirrels (GC treated = 232 rattles from 10 squirrels, positive control = 367 rattles from 12 squirrels, and negative control = 115 rattles from 23 squirrels). Based on a cross-validated assessment of the accuracy of our approach (see details in Siracusa et al. forthcoming), 52% of all focal rattles should have been identified correctly as focal rattles (i.e., 48% incorrectly classified as coming from a neighbor and, therefore, excluded; false negative error rate = 48%) and 6% of the rattles labeled as focal rattles (after manually removing the nonrattles) should actually have been neighbor rattles (i.e., false error rate of 6%). Therefore, although our final data set included only half of all rattles produced by our focal squirrels during their 24-h trials, the vast majority of rattles that were included in the data set were from the focal individual.

Acoustic analysis

We used Avisoft SASLab Pro software version 5.0 (Avisoft, 2015) to analyze the acoustic structure of rattles recorded in both experiments. The rattles were loaded into Avisoft, and for each rattle we generated a spectrogram (Fast Fourier Transform size: 512, window: hamming, temporal resolution: 1.45 ms, frequency resolution: 86 Hz, overlap: 87.5%) and the program extracted the acoustic parameters of interest (described below) using an existing protocol for rattle acoustic analysis. We oversaw this process manually, checking that each call was recognized and analyzed in its entirety by AviSoft, and that none were cutoff—if the program did not recognize the call in its entirety, we would adjust the recognition parameters slightly. To characterize rattles, we measured 3 acoustic parameters: rattle duration, mean frequency (the frequency below which lies 50% of the energy of the signal, as measured from an averaged power spectrum of the entire signal), and entropy, a measure of noisiness of a signal. Because rattles are broadband and noisy signals, meaning that the majority of the energy in a call is dispersed across the frequency domain, mean frequency is a more appropriate measure of the frequency of the call than peak frequency. AviSoft measures Weiner Entropy (spectral flatness), calculated by dividing the geometric mean of the power spectrum by the arithmetic mean of the power spectrum, which ranges from 0 (pure tone) to 1 (white noise). We limited analysis to these 3 acoustic variables because in our review of the literature, these variables appeared to be most commonly influenced by arousal (Manser 2001; Rendall 2003; Facchini et al. 2005; Soltis et al. 2005b; Slocombe et al. 2009; Esch 2009; Zimmerman 2009). These

measurements were made using the “automatic parameter measurements” feature of SASLab Pro to eliminate human bias in the measurements (settings: threshold -13 dB, hold time of 150 ms).

Because high frequencies attenuate more readily than low frequencies, entropy and mean frequency could, in theory, covary with recording distance. In the capture-induced stress experiment, a constant recording distance of approximately 5 m was maintained for all recordings. In the GC-induced stress experiment, in which rattles were recorded on stationary zoom microphones, to ensure that recording distance did not vary with time or treatment, we measured the signal-to-noise ratio of a subset of 140 rattles and found no significant relationships between rattle amplitude (a proxy for recording distance) and time of day (linear regression: $t = -1.33$, degrees of freedom [df] = 6.6, $P = 0.19$) or treatment (linear regression: $t = -1.66$, df = 24.9, $P = 0.11$). This indicates that any variation in rattle entropy throughout the day or among the treatments was not due to focal squirrels being closer to or further from the microphone.

Statistical analyses

For statistical analyses, we used R (version 3.5.1; R Developmental Core Team 2018) with the package lme4 (version 1.17; Bates et al. 2015) to fit linear mixed-effects models and lmerTest version 3.0 (Kuztsova et al. 2017) to assess the significance of these models. For the capture-induced stress experiment, we included rattle-collection method (post-trap or opportunistic) as a fixed effect. We included squirrel ID as a random effect because we analyzed multiple rattles from the same squirrels across multiple years.

Wilson et al. (2015) found no effects of age, sex, or Julian date on the acoustic structure of rattles recorded from this same population. To confirm this finding, we ran separate linear mixed-effects models with each variable included as a fixed effect and found no significant relationship between any of these variables and any of the 3 acoustic structural features. Age had no relationship with acoustic structure in either stressed (*duration*: $t = -0.08$, df = 36.9, $P = 0.94$; *mean frequency*: $t = -0.54$, df = 34.5, $P = 0.59$; *entropy*: $t = -1.26$, df = 36.9, $P = 0.21$) or unstressed (*duration*: $t = -0.04$, df = 51.5, $P = 0.97$; *mean frequency*: $t = -0.66$, df = 152.14, $P = 0.51$; *entropy*: $t = -1.01$, df = 138.4, $P = 0.31$) squirrels. The same was true for sex—neither stressed (*duration*: $t = <0.001$, df = 36.81, $P = 0.99$; *mean frequency*: $t = 0.48$, df = 33.7, $P = 0.96$; *entropy*: $t = 0.34$, df = 36.6, $P = 0.74$) nor unstressed (*duration*: $t = 0.93$, df = 162.2, $P = 0.36$; *mean frequency*: $t = 0.08$, df = 154.7, $P = 0.94$; *entropy*: $t = -0.91$, df = 176.5, $P = 0.36$) squirrels showed any such relationship. And the same was true for Julian date, in both stressed (*duration*: $t = -0.05$, df = 34.55, $P = 0.81$; *mean frequency*: $t = 0.78$, df = 35.11, $P = 0.55$; *entropy*: $t = 1.05$, df = 34.23, $P = 0.59$) and unstressed squirrels (*duration*: $t = 0.10$, df = 161.22, $P = 0.81$; *mean frequency*: $t = -0.03$, df = 162.33, $P = 0.37$; *entropy*: $t = 0.85$, df = 161.58, $P = 0.45$).

We also found no year effects for any of the acoustic parameters measured—we conducted linear mixed models for each acoustic variable and found no effect of year on any variable in both stressed (*duration*: $F_{2, 36.0} = 0.71$, $P = 0.50$; *mean frequency*: $F_{2, 36.2} = 2.20$, $P = 0.08$; *entropy*: $F_{2, 37.1} = 3.20$, $P = 0.10$) and unstressed (*duration*: $F_{3, 260.5} = 1.76$, $P = 0.16$; *mean frequency*: $F_{3, 261.8} = 2.58$, $P = 0.09$; *entropy*: $F_{3, 262.3} = 2.21$, $P = 0.09$) squirrels.

To examine the effects of administration of exogenous GCs on the acoustic structure of rattles, we fit 3 separate linear mixed-effects models—one for each of the 3 acoustic response variables (duration, mean frequency, entropy). Each model included an interaction

between treatment group and time since treatment consumption (both linear and quadratic terms) as fixed effects and squirrel ID ($n = 44$) as a random effect. In order to include the rattles of negative control group squirrels in this model, we found the average time at which the GC-treated and positive control (fed) squirrels consumed their treatment (1015 h) and set that as time of treatment consumption for all negative control group (unfed) squirrels. For example, a rattle emitted at 1030 h would have a “time since treatment” value of 900 s and a rattle emitted at 1000 h would have a time since treatment value of -900 s. Time since treatment consumption was standardized (mean [time of day] = 0, standard deviation [SD] = 1). In both experiments, Q-Q plots were generated to test for normality, and residuals were found to be normally distributed.

We did not compare the rattles of GC-treated squirrels directly with the rattles of trap-stressed squirrels for two primary reasons. The first is that for the GC-treated squirrels, because we checked for consumption of treatment only every hour, we have far less precision in determining which rattles occurred at peak circulating GC levels (90–120 min post-treatment). Thus, it was impossible to compare the post-trap rattles produced at peak stress levels with rattles produced at peak GC levels. The second reason is that the rattles in the post-trap stress experiment were recorded with a different, slightly higher quality recording unit than those recorded in the GC treatment experiment, making direct comparisons across recordings inappropriate.

If elevated plasma GCs alter rattle acoustic structure, we expected that the effects of the GC treatment on rattle acoustic structure would be strongest within 90–120 min of treatment consumption, the time frame in which circulating GCs should be highest using this treatment paradigm (Breuner et al. 1998; van Kesteren et al. 2018 preprint). Thus, we included a nonlinear (quadratic) term for time since treatment consumption and its interaction with treatment because we expected that the effects of the treatment would exhibit a nonlinear relationship, peaking within 90–120 min of treatment and then declining throughout the remainder of the day.

RESULTS

Effects of capture-induced stress on rattle acoustic structure

Capture-induced stress caused pronounced differences in rattle acoustic structure: post-trap rattles were longer, higher in frequency, and noisier than rattles collected opportunistically. Thus, there was indeed a clear stress signature. The average duration of post-trap rattles (4.77 ± 2.25 (SD) s) was significantly longer than that of opportunistic rattles (2.93 ± 1.28 s), a 63% increase ($t = 3.78$, df = 209.41, $P < 0.001$; Figure 1A). The average mean frequency of post-trap rattles (7269.53 ± 1180.76 Hz) was slightly but significantly higher than that of opportunistic rattles (6971.753 ± 1007.37 Hz), a 4.3% increase ($t = 2.82$, df = 218.01, $P = 0.005$; Figure 1B). And finally, the average entropy of post-trap rattles (0.754 ± 0.035) was slightly but significantly higher than that of opportunistic rattles (0.712 ± 0.047), a 5.9% increase ($t = 4.14$, df = 78.52, $P < 0.001$; Figure 1C).

Effects of administration of GCs on rattle acoustic structure

Administration of exogenous GCs did not produce the same effects on rattle acoustic structure as capture-induced stress—the rattle

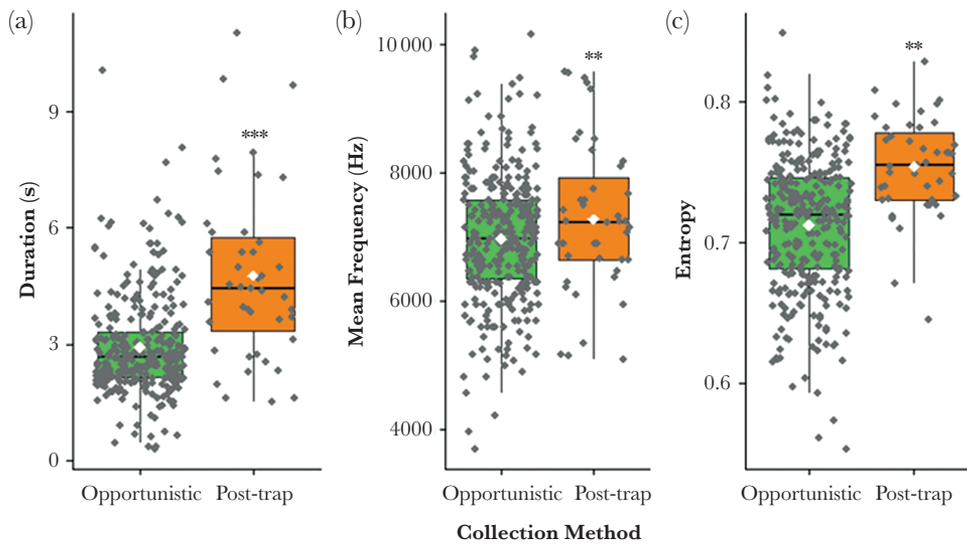


Figure 1 Effects of short-term stress (live-trapping and handling) on rattle (A) duration (s), (B) mean frequency (Hz), and entropy. Post-trap rattles were collected within a minute of the squirrel exiting a trap and rattles collected opportunistically were collected from unprovoked squirrels. Post-trap rattles were significantly longer, higher in frequency, and higher in entropy. The black lines denote median, the white diamonds denote mean. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$.

Table 2 Compared with the rattles of GC-treated squirrels, the duration of rattles of negative control squirrels increases linearly as a function of time since consumption of treatment

Fixed effect	<i>b</i>	Standard error (SE)	df	<i>t</i>	<i>P</i> -value
Intercept	2.63	0.34	0.74	7.64	0.14
Time since treatment	0.07	0.07	683.0	0.90	0.37
Positive control treatment	0.07	0.21	31.35	0.32	0.75
Negative control treatment	0.32	0.44	0.9	0.72	0.61
Time since treatment ²	-0.02	0.08	683.7	-0.30	0.76
Time since treatment × positive control	0.07	0.09	681.04	0.73	0.47
Time since treatment × negative control	0.33	0.12	692.2	2.67	0.01
Time since treatment ² × positive control	-0.06	0.09	686.4	-0.68	0.50
Time since treatment ² × negative control	0.17	0.13	701.4	1.29	0.20

No other effects of treatment were found. Output shown is from a linear mixed-effects model that assessed the influence of administration of GCs on rattle duration compared with those fed supplemental food (positive control) or unfed squirrels (negative control), including time since treatment as both a linear and quadratic term. Individual identity was included as a random effect. GC treatment is in the intercept. Sample size: 714 rattles (GC treated: $n = 232$, positive control: $n = 367$, negative control: $n = 115$). Bolded terms are significant.

acoustic features of GC-treated squirrels did not follow the predicted pattern of peaking after treatment and then declining as a function of time since treatment (Tables 2–4, Figure 2). There was, however, a significant linear interaction between treatment and the amount of time elapsed since treatment consumption on rattle duration ($F_{2,677.4} = 3.78, P = 0.02$). This effect was largely driven by the increases in rattle duration observed in negative control group squirrels (Figure 2A): rattles from negative control group squirrels increased in length throughout the day compared with those treated with GCs ($b = 0.33, t = 2.67, P = 0.01$, Table 2, Figure 2A). Rattle durations of squirrels treated with GCs did not change differentially over the course of the day when compared with rattle durations of squirrels fed peanut butter only (positive control group: $b = 0.07, t = 0.73, P = 0.47$, Table 2, Figure 2A).

There were no treatment effects on rattle mean frequency ($F_{2,56} = 0.60, P = 0.63$, Table 3) or entropy ($F_{2,56} = 0.47, P = 0.63$, Table 4) and the effects of the treatments on rattle mean frequency

or entropy did not depend on the amount of time that had elapsed since treatment consumption, as indicated by the lack of interactions between treatment and time elapsed since treatment consumption (both linear and quadratic terms). However, the mean frequency of rattles from squirrels recorded in all 3 treatment groups increased throughout the day ($F_{1,683.3} = 4.77, P = 0.03$). Overall, there were no significant nonlinear effects of time since treatment consumption or its interaction with treatment on rattle duration, frequency, or entropy (Tables 2–4).

DISCUSSION

Our study shows that short-term stress, in this case induced by live capture and handling, significantly influences the acoustic structure of territorial vocalizations in red squirrels. Squirrels experiencing capture-induced stress produced rattles that were longer in duration, higher in frequency, and noisier (higher entropy) than rattles

Table 3

No effects of treatment on rattle mean frequency were found; however, mean frequency did increase as a function of time since treatment consumption

Fixed effect	<i>b</i>	SE	df	<i>t</i>	<i>P</i> -value
Intercept	9184.42	211.67	0.52	43.4	0.09
Time since treatment	175.50	77.15	681.7	2.28	0.02
Positive control treatment	5.73	281.57	43.15	0.02	0.98
Negative control treatment	-268.48	291.95	0.7	-0.92	0.57
Time since treatment ²	-75.89	78.46	682.3	-0.97	0.33
Time since treatment × positive control	-147.53	95.56	680.7	-1.54	0.12
Time since treatment × negative control	-64.04	129.60	693.1	-0.49	0.62
Time since treatment ² × positive control	-114.80	95.99	683.4	-1.20	0.23
Time since treatment ² × negative control	21.40	136.18	677.0	0.16	0.88

Output shown is from a linear mixed-effects model that assessed the influence of administration of GCs on rattle mean frequency (Hz) compared with those fed supplemental food (positive control) or unfed squirrels (negative control), including time since treatment as both a linear and quadratic term. Individual identity was included as a random effect. Sample size: 714 rattles (GC treated: *n* = 232, positive control: *n* = 367, negative control: *n* = 115). Bolded terms are significant.

Table 4

No effects of treatment were found on rattle entropy

Fixed effect	<i>b</i>	SE	df	<i>t</i>	<i>P</i> -value
Intercept	7.86e-1	9.27e-3	4.70e+1	84.82	<2e-16
Time since treatment	-8.09e-4	3.14e-3	6.80e+2	-0.26	0.80
Positive control treatment	-4.03e-3	1.26e-2	4.36e+1	-0.32	0.75
Negative control treatment	7.49e-3	1.26e-2	6.63e+1	0.59	0.56
Time since treatment ²	1.06e-3	3.20e-3	6.81e+2	0.33	0.74
Time since treatment × positive control	6.11e-4	3.90e-3	6.79e+2	0.16	0.88
Time since treatment × negative control	3.86e-3	5.31e-3	7.04e+2	0.73	0.47
Time since treatment ² × positive control	-3.09e-3	3.91e-3	6.81e+2	-0.79	0.43
Time since treatment ² × negative control	-1.10e-4	5.56e-3	7.02e+2	-0.02	0.98

Output shown is from a linear mixed-effects model that assessed the influence of administration of GCs on rattle entropy compared with those fed supplemental food (positive control) or unfed squirrels (negative control), including time since treatment as both a linear and quadratic term. Individual identity was included as a random effect. GC treatment is in the intercept. Bolded terms are significant. Sample size: 714 rattles (GC treated: *n* = 232, positive control: *n* = 367, negative control: *n* = 115). Bolded terms are significant.

produced by positive control squirrels. However, we were unable to reproduce these same effects on acoustic structure by experimentally increasing circulating GCs, despite the fact that GCs increase in response to trapping and handling (Bossion et al. 2012; van Kesteren et al. 2018 preprint). Indeed, the rattles of squirrels treated with GCs did not exhibit the expected structural distinctions from the rattles of positive control or negative control squirrels over the treatment period.

The only significant differences in rattle acoustic structure between treatment groups in the GC treatment experiment was that in negative control squirrels, rattles increased in duration throughout the day, whereas the rattles of GC-treated and control (fed) squirrels did not significantly vary throughout the day. This may indicate that supplemental feeding suppresses a normally occurring pattern in rattle structure—rattles may normally increase in duration throughout the day. Differences in study grids may also explain this difference: squirrels in the GC-treated and positive control groups were on a different study grid than those in the negative control group. The study grid on which the GC-treated and positive control squirrels lived has historically had higher squirrel population densities than the study grids on which the negative control squirrels lived; it is thus possible that at lower population densities, squirrels produce longer rattles. Both of these possible explanations, however, would need to be examined directly.

The effects of short-term stress (trapping and handling) on rattle acoustic structure that we observed (longer duration, higher mean frequency, and higher entropy) are largely congruent with such trends in acoustic structure in relation to stress in many species. Chimpanzee screams, for example, increase in duration with the severity of an agonistic encounter (Slocombe et al. 2009). In dog barks (*Canis lupus familiaris*, Tokuda et al. 2002), human infant cries (Facchini et al. 2005), baboon grunts (*Papio hamadrayas*, Rendall 2003), and meerkat alarm calls (*Suricata suricatta*, Manser 2001), noisiness (entropy) increases with short-term stress. In many species, an increase in short-term stress is associated with an increase in pitch-related characteristics. For example, during capture–release events, female bottlenose dolphins with dependent calves produce whistles of elevated frequency (*Tursiops truncatus*, Esch 2009). The same pattern is observed in adult female African elephants (*Loxondota africana*, Soltis et al. 2005b), tree shrews (*Tupaia belangeri*, Schehka and Zimmermann 2009), and zebra finches (Perez et al. 2012): short-term stress is associated with an increase in vocalization pitch. In giant panda cubs (*Ailuropoda melanoleuca*), increased stress is associated with all of the trends in acoustic structure that we observed in post-trap rattles: longer duration, higher frequency, and increased noise (Stoeger et al. 2012).

Our results somewhat resemble those of Perez et al. (2012), who investigated how an environmental stressor (social isolation) and

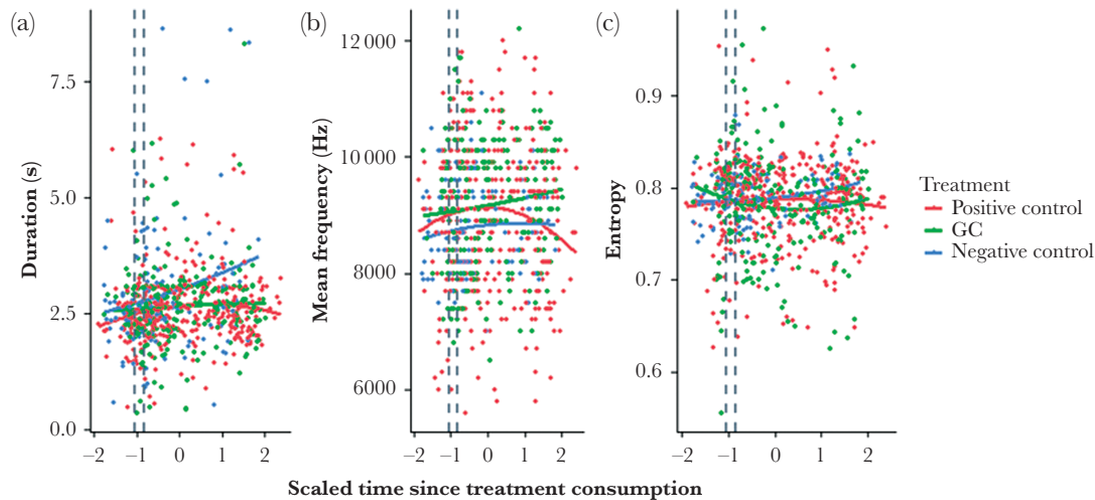


Figure 2

Effects of exogenous GCs (“GC”) and supplemental food (“Positive control”) on rattle (A) duration (s), (B) mean frequency (Hz), and (C) entropy as a function of time since treatment. For negative control squirrels, “time of treatment” is standardized at 1015 h, the average time of treatment consumption of GC-treated and control (fed) squirrels. The vertical gray dotted lines indicate the 1-h time frame in which squirrels consumed their treatments. Squirrels fed supplemental food (positive controls), exogenous GCs, or those that were negative controls had similar acoustic structure except that negative control squirrels had significantly longer rattles than GC-treated squirrels as the time since treatment consumption increased (Table 2). Time since treatment consumption was standardized (mean = 0, SD = 1).

treatment with exogenous GCs affected vocalization structure in zebra finches. In their study, social isolation induced vocalizations of increased duration and pitch and reduced overall vocal activity. However, oral administration of GCs only resulted in vocalizations with increased pitch, but no other effects were observed (Perez et al. 2012). The results from Perez et al. (2012) and our study suggest that short-term stressors alter vocalization structure but any increases in GCs caused by the short-term stressor are not solely responsible for these changes.

Our findings and those of Perez et al. (2012) suggest that the acoustic structure of vocalizations can be altered by short-term stress, but the relationship between circulating GC levels and acoustic structure of vocalizations is not straightforward. GC treatment and capture-induced stress result in comparable concentrations of plasma GCs (van Kesteren et al. 2018 preprint), indicating that our GC treatment regime fairly accurately simulates the increase in plasma GCs experienced as a result of capture. Thus, other hormones or neurochemicals may be implicated in modulation of the acoustic structure of vocalizations. For example, in rat pups, several classes of dopamine receptor agonists reduced the production of stress-induced ultrasonic vocalizations caused by isolation; this is a sign of reduced separation anxiety (Dastur et al. 1999). It is also possible that the acoustic structure of vocalizations has a nonmonotonic dose–response relationship with GCs. There is precedent for such a relationship: in white crowned sparrows, moderate doses of corticosterone induced elevated physical activity, whereas high levels did not (Breuner et al. 1998). We only provisioned squirrels with one dosage of GCs and so were unable to address whether lower or higher dosages of GCs would alter rattle acoustic structure. Together, this suggests the importance of considering additional mechanisms that may underlie the observed changes in vocalization acoustic structure.

Because treatment with exogenous GCs induced none of the expected changes to rattle acoustic structure, it is possible that the acoustic changes observed in the rattles of trapped squirrels

were produced by an effect of trapping besides increases in GCs. Because rattles function to advertise territorial ownership, it is possible that a squirrel that has been in a trap and unable to defend its territory for up to 2 h, on release, compensates by producing rattles that are longer and noisier. This hypothesis, however, would require explicit tests.

It is also worth considering here the possible influence of energetic state on rattle acoustic structure. The 2 manipulated groups (GC and positive control) were provisioned with 10 g of a peanut butter/wheat germ mixture, and it is conceivable that this energetic boost impacted rattle structure. A red squirrel’s daily energetic expenditure ranges from approximately 177 to 660 kJ/day, depending on season and reproductive state; during mid-summer, when our experiment was conducted, a male squirrel’s daily energetic expenditure likely falls near the middle of that range (Fletcher et al. 2012). Ten grams of natural peanut butter and wheat germ contains approximately 240 kJ; thus, peanut butter treatments constituted a significant portion of a squirrel’s daily energy requirements. A significant body of literature has examined the effects of energetic state on vocalizations. Most notably, studies on songbirds have found that birds provisioned with supplemental food in the evening produce longer and more complex dawn choruses the following morning (Cuthill and MacDonald 1990; Berg et al. 2005; Barnett and Briskie 2007).

It is thus conceivable that the energetic boost provided by peanut butter treatments interfered with any effects of GCs on acoustic structure. However, given the lack of significant differences in the rattles of squirrels in any of the 3 treatment groups, we consider this possibility unlikely.

Our findings constitute further evidence that territorial vocalizations such as rattles contain more information than territorial ownership. In red squirrels, rattles have not only the capacity to communicate stable information about the signaler’s individual identity and potential kin relationships (Digweed et al. 2012; Wilson et al. 2015; Shonfield et al. 2017) but also labile information, such

as short-term stress. In some cases, it is possible that this stable and labile information may interact—the stress state of the signaler might modify the ability of conspecifics to discriminate whether they are kin or nonkin, as proposed by Shonfield et al. (2017). This layering of stable and labile encoded information in vocalizations may not be uncommon, appearing across a number of animal taxa (Rendall 2003; Seyfarth and Cheney 2003; Blumstein and Munos 2005; Soltis 2005a; Koren and Geffen 2009; Terleph et al. 2016).

There are several hypotheses on the functional significance of these tendencies in vocalizations associated with high-stress contexts. In social species, the unpredictability hypothesis states that calls that contain more nonlinearities are more difficult to habituate to, and thus noisy alarm calls are more likely to capture the attention of a conspecific in the event of a predatory or otherwise dangerous event (Fitch et al. 2002, Blumstein and Recapet 2009). Another hypothesis holds that screams produced when an animal is in imminent danger of predation serve to either startle and distract the predator, or solicit intervention from another animal, either a social group member or a “pirate” predator that may attempt to steal the prey and unintentionally free it (Hogstedt 1982). In the case of red squirrels, one hypothesis that can be envisaged is that honestly communicating stress to neighbors may advertise a willingness to aggressively defend one’s territory. Another possibility is that instead of honestly depicting a willingness to defend a territory, vocal cues of stress might inadvertently reveal that the caller faces some other challenge and might, therefore, be less capable of defending their territory. These two hypotheses, however, would need to be tested directly—for example, a playback study could test whether the rattles of stressed squirrels are more or less likely to deter territorial intrusions from neighboring squirrels than rattles of unstressed squirrels. If stress-influenced rattles are more likely to deter intruders, and if their production predicts an attack or further escalation by the signaler, then stressed rattles would be considered aggressive signals (Searcy and Beecher 2009); if the opposite was the case, they would be considered index signals (Smith and Harper 1995).

Though research on stress-induced changes to vocalizations has focused primarily on group-living species, the encoding of labile information such as short-term stress in vocalizations may have consequences in a population of solitary, territorial animals as well, perhaps enabling neighbors to eavesdrop on the physiological state of the signaler and adjust their own behavior or reproduction accordingly. Eavesdropping by conspecifics, or the acquisition of public information, may have important ecological consequences (Valone 2007; Schmidt et al. 2010). For example, in many species, including red squirrels (Fisher et al. 2017; Lane et al. 2018), breeding earlier than other individuals in your population may be advantageous. Cues about the physiological state of a signaler contained in territorial vocalizations may provide an important source of information about when other individuals in the population are breeding—in red squirrels, the strongest level of selection for post-natal growth rate and birth date is the social neighborhood (Fisher et al. 2017). As such, labile information contained in vocalizations, such as stress state, may have broader ecological consequences by serving as public information and modifying the timing of reproduction in seasonally breeding species.

Overall, our results indicate that red squirrel territorial vocalizations may contain labile information on physiological state, in addition to the previously documented stable information about territorial ownership and individual identity. This study represents one of only very few experimental tests of effects of stress on acoustic structure in any species. Future studies could explore the

possible interactions between stable and labile information encoded in these calls, and the ways in which these layers of encoded information might influence behavioral or reproductive dynamics.

FUNDING

Funding was provided by the University of Michigan and National Science Foundation (IOS 1749627 to B.D.) as well as the Natural Sciences and Engineering Council of Canada (to S.B., M.M.H., J.E.L., A.G.M., and D.W.).

We thank the Champagne and Aishihik First Nations for providing access to the land on which the study sites for this project were located, in particular Agnes MacDonald and her family for long-term access to her trapline. We also thank Zach Fogel and Noah Israel, the diligent field technicians whose work on the GC-induced experiment was crucial for its success. This is publication 99 of the Kluane Red Squirrel Project.

Data accessibility: Analyses reported in this article can be reproduced using the data provided by Sehrsweeney et al. (2019).

Handling editor: Amanda Ridley

REFERENCES

- Andersson M. 1994. Sexual selection. Princeton (NJ): Princeton University Press.
- August PV, Anderson JGT. 1987. Mammal sounds and motivation-structural rules: a test of the hypothesis. *J Mammal*. 68:1–9.
- Barnett CA, Briskie JV. 2007. Energetic state and the performance of dawn chorus in silvereyes (*Zosterops lateralis*). *Behav Ecol Sociobiol*. 61:579–587.
- Bates D, Maechler M, Bolker B, Walker S. 2015. Fitting linear mixed-effects models using lme4. *J Stat Soft*. 67:1–48.
- Bayart F, Hayashi KT, Faull KF, Barchas JD, Levine S. 1990. Influence of maternal proximity on behavioral and physiological responses to separation in infant rhesus monkeys (*Macaca mulatta*). *Behav Neurosci*. 104:98–107.
- Bee MA, Perrill SA, Owen PC. 1999. Size assessment in simulated territorial encounters between male green frogs (*Rana clamitans*). *Behav Ecol Sociobiol*. 45:177–184.
- Beecher MD. 1989. Signaling systems for individual recognition: an information theory approach. *Anim Behav*. 38:248–261.
- Beer CG. 1970. On the responses of laughing gull chicks (*Larus atricilla*) to the calls of adults II. Age changes and responses to different types of call. *Anim Behav*. 18:661–677.
- Berg ML, Beintema NH, Welbergen JA, Komdeur J. 2005. Singing as a handicap: the effects of food availability and weather on song output in the Australian Reed Warbler *Acrocephalus australis*. *Oikos*. 36:102–109.
- Biben M, Symes D, Masataka N. 1986. Temporal and structural analysis of affiliative vocal exchanges in squirrel monkeys (*Saimiri sciureus*). *Behaviour*. 98:259–273.
- Blumstein D, Richardson D, Cooley L, Winternitz J, Daniel J. 2008. The structure, meaning and function of yellow-bellied marmot pup screams. *Anim Behav*. 76:1055–1064.
- Blumstein DT, Munos O. 2005. Individual, age and sex-specific information is contained in yellow-bellied marmot alarm calls. *Anim Behav*. 69:353–361.
- Blumstein DT, Récapet C. 2009. The sound of arousal: the addition of novel non-linearities increases responsiveness in marmot alarm calls. *Ethol*. 115:1074–1081.
- Bosson CO, Islam Z, Boonstra R. 2012. The impact of live trapping and trap model on the stress profiles of North American red squirrels. *J Zool*. 288:159–169.
- Breuner CW, Greenberg AL, Wingfield JC. 1998. Noninvasive corticosterone treatment rapidly increases activity in Gambel’s white-crowned sparrows (*Zonotrichia leucophrys gambelii*). *Gen Comp Endocrinol*. 111:386–394.
- Clark AP. 1993. Rank differences in the production of vocalizations by wild chimpanzees as a function of social context. *Am J Primatol*. 31:159–197.
- Clutton-Brock TH, Albon SD. 1979. Roaring of red deer and evolution of honest advertisement. *Behaviour*. 69:145–170.

- Cuthill IC, Macdonald WA. 1990. Experimental manipulation of the dawn and dusk chorus in the blackbird (*Turdus merula*). *Behav Ecol Sociobiol*. 26:209–216.
- Dantzer B, Newman AE, Boonstra R, Palme R, Boutin S, Humphries MM, McAdam AG. 2013. Density triggers maternal hormones that increase adaptive offspring growth in a wild mammal. *Science*. 340:1215–1217.
- Dastur FN, McGregor IS, Brown RE. 1999. Dopaminergic modulation of rat pup ultrasonic vocalizations. *Eur J Pharmacol*. 382:53–67.
- Digweed S, Rendall D, Imbeau T. 2012. Who's your neighbor? Acoustic cues to individual identity in red squirrel (*Tamiasciurus hudsonicus*) rattle calls. *Curr Zool*. 58:758–764.
- Digweed SM, Fedigan LM, Rendall D. 2007. Who cares who calls? Selective responses to the lost calls of socially dominant group members in the white-faced capuchin (*Cebus Capucinus*). *Am J Primatol*. 69:829–835.
- Donald JL, Boutin S, Steele MA. 2011. Intraspecific cache pilferage by larder-hoarding red squirrels (*Tamiasciurus hudsonicus*). *J Mammal*. 92:1013–1020.
- Ehret G. 2005. Infant rodent ultrasounds – a gate to the understanding of sound communication. *Behav Genet*. 35:19–29.
- Esch HC, Sayigh LS, Blum JE, Wells RS. 2009. Whistles as potential indicators of stress in bottlenose dolphins (*Tursiops truncatus*). *J Mammal*. 90:638–650.
- Ey E, Hammerschmidt K, Seyfarth RM, Fischer J. 2007. Age and sex-related variations in clear calls of *Papio ursinus*. *Int J Primatol*. 28:947–960.
- Facchini A, Bellieni CV, Marchettini N, Pulselli FM, Tiezi EBP. 2005. Relating pain intensity of newborns to onset of nonlinear phenomena in cry recordings. *Phys Lett A*. 338:332–337.
- Fisher DN, Boutin S, Dantzer B, Humphries MM, Lane JE, McAdam AG. 2017. Multilevel and sex-specific selection on competitive traits in North American red squirrels. *Evolution*. 71:1841–1854.
- Fitch WT. 1997. Vocal tract length and formant frequency dispersion correlate with body size in rhesus macaques. *J Acoust Soc Am*. 102:1213–1222.
- Fitch WT, Neubauer J, Herzel H. 2002. Calls out of chaos: the adaptive significance of nonlinear phenomena in mammalian vocal production. *Anim Behav*. 63:407–418.
- Fletcher QE, Landry-Cuerrier M, Boutin S, McAdam AG, Speakman JR, Humphries MM. 2013. Reproductive timing and reliance on hoarded capital resources by lactating red squirrels. *Oecologia*. 173:1203–1215.
- Fletcher QE, Speakman JR, Boutin S, McAdam AG, Sarah B, Humphries MM. 2012. Seasonal stage differences overwhelm environmental and individual factors as determinants of energy expenditure in free-ranging red squirrels. *Funct Ecol*. 26:277–287.
- Hogstedt G. 1982. Adaptation unto death: function of fear screams. *Am Nat*. 121:562–570.
- Koren L, Geffen E. 2009. Complex call in male rock hyrax (*Procapra capensis*): a multi-information distributing channel. *Behav Ecol Sociobiol*. 63:581–590.
- Koren L, Mokady O, Geffen E. 2008. Social status and cortisol levels in singing rock hyraxes. *Horm Behav*. 54:212–216.
- Krebs CJ, Boutin SA, Boonstra R. 2001. Ecosystem dynamics of the boreal forest: the Kluane project. New York (NY): Oxford University Press.
- Kuznetsova A, Brockhoff P, Christianson HB. 2017. lmerTest package: tests in linear mixed effects models. *J Stat Soft*. 82:1–26.
- Lane JE, McAdam AG, McFarlane SE, Williams CT, Humphries MM, Coltman DW, Gorrell JC, Boutin S. 2018. Phenological shifts in North American red squirrels: disentangling the roles of phenotypic plasticity and microevolution. *J Evol Biol*. 31:810–821.
- Larsen KW, Boutin S. 1994. Movements, survival, and settlement of red squirrel (*Tamiasciurus hudsonicus*) offspring. *Ecology*. 75:214–223.
- Lingle S, Rendall D, Pellis SM. 2007. Altruism and recognition in the anti-predator defense of deer: 1. Species and individual variation in fawn distress calls. *Anim Behav*. 73:897–905.
- Manser MB. 2001. The acoustic structure of suricates' alarm calls varies with predator type and the level of response urgency. *Proc Biol Sci*. 268:2315–2324.
- Masataka N, Symmes D. 1986. Effect of separation distance on isolation call structure in squirrel monkeys (*Saimiri sciureus*). *Am J Primatol*. 10:271–278.
- McAdam A, Boutin S, Sykes A, Humphries M. 2007. Life histories of female red squirrels and their contributions to population growth and lifetime fitness. *Ecoscience*. 14:362–369.
- Morton ES. 1977. On the occurrence and significance of motivation-structural rules in some bird and mammal sounds. *Am Nat*. 111:855–869.
- Muller MN, Wrangham RW. 2004. Dominance, aggression and testosterone in wild chimpanzees: a test of the “challenge hypothesis.” *Anim Behav*. 67:113–123.
- Perez EC, Elie JE, Soulage CO, Soula HA, Mathevon N, Vignal C. 2012. The acoustic expression of stress in a songbird: does corticosterone drive isolation-induced modifications of zebra finch calls? *Horm Behav*. 61:573–581.
- Perez EC, Mariette MM, Cochard P, Soulage CO. 2016. Corticosterone triggers high-pitched nestlings' begging calls and affects parental behavior in the wild zebra finch. *Behav Ecol*. 27:1665–1677.
- R Core Team. 2018. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing.
- Reby D, McComb K. 2003. Anatomical constraints generate honesty: acoustic cues to age and weight in the roars of red deer stags. *Anim Behav*. 65:519–530.
- Rendall D. 2003. Acoustic correlates of caller identity and affect intensity in the vowel-like grunt vocalizations of baboons. *J Acoust Soc Am*. 113:3390–3402.
- Sacchi R, Saino N, Galeotti P. 2002. Features of begging calls reveal general condition and need of food of barn swallow (*Hirundo rustica*) nestlings. *Behav Ecol*. 13:268.
- Sapolsky RM, Romero LM, Munck AU. 2000. How do glucocorticoids influence stress responses? Integrating permissive, suppressive, stimulatory, and preparative actions. *Endocr Rev*. 21:55–89.
- Schehka S, Zimmermann E. 2009. Acoustic features to arousal and identity in disturbance calls of tree shrews (*Tupaia belangeri*). *Behav Brain Res*. 203:223–231.
- Schmidt K, Dall S, Van Gils J. 2010. The ecology of information: an overview on the ecological significance of making informed decisions. *Oikos*. 119:304–316.
- Searcy WA, Beecher MD. 2009. Song as an aggressive signal in songbirds. *Anim Behav*. 78:1281–1292.
- Schrsweaney M, Wilson DR, Bain M, Boutin S, Lane JE, McAdam AG, Dantzer B. 2019. Data from: the effects of stress and glucocorticoids on vocalizations: a test in North American red squirrels. Dryad Digital Repository. doi:10.5061/dryad.9c6k203.
- Seyfarth RM, Cheney DL. 2003. Signalers and receivers in animal communication. *Annu Rev Psychol*. 54:145–173.
- Shonfield J, Gorrell JC, Coltman DW, Boutin S. 2017. Using playback of territorial calls to investigate mechanisms of kin discrimination in red squirrels. *Behav Ecol*. 28:1–9.
- Siracusa E, Morandini M, Boutin S, Humphries MM, Dantzer B, Lane JE, McAdam AG. 2017. Red squirrel territorial vocalizations deter intrusions by conspecific rivals. *Behaviour*. 154:1259–1273.
- Siracusa ER, Wilson DR, Studd EK, Boutin S, Humphries MM, Dantzer B, Lane JE, McAdam AG. 2019. North American red squirrels mitigate costs of territory defence through social plasticity. *Anim Behav*. doi:10.1016/j.anbehav.2019.02.014
- Slocombe KE, Townsend SW, Zuberbühler K. 2009. Wild chimpanzees (*Pan troglodytes schweinfurthii*) distinguish between different scream types: evidence from a playback study. *Anim Cogn*. 12:441–449.
- Smith CC. 1968. The adaptive nature of social organization in the genus of three squirrels *Tamiasciurus*. *Ecol Monograph*. 38:31–64.
- Smith CC. 1978. Structure and function of the vocalizations of tree squirrels (*Tamiasciurus hudsonicus*). *J Mammal*. 59:793–808.
- Smith MJ, Harper DCG. 1995. Animal signals: models and terminology. *J Theor Biol*. 177:305–311.
- Soltis J, Leong K, Savage A. 2005a. African elephant vocal communication I: antiphonal calling behaviour among affiliated females. *Anim Behav*. 70:579–587.
- Soltis J, Leong K, Savage A. 2005b. African elephant vocal communication II: rumble variation reflects the individual identity and emotional state of callers. *Anim Behav*. 70:589–599.
- Stoeger AS, Baotic A, Li D, Charlton BD. 2012. Acoustic features indicate arousal in infant giant panda vocalisations. *Ethol*. 118:896–905.
- Sueur J, Aubin T, Simonis C. 2008. Seewave, a free modular tool for sound analysis and synthesis. *Bioacoustics*. 18:213–226.
- Terleph TA, Malaivijitnond S, Reichard UH. 2016. Age related decline in female lar gibbon great call performance suggests that call features correlate with physical condition. *BMC Evol Biol*. 16:4.

- Tokuda I, Riede T, Neubauer J, Owren MJ, Herzl H. 2002. Nonlinear analysis of irregular animal vocalizations. *J Acoust Soc Am.* 111:2908–2919.
- Valone T. 2007. From eavesdropping on performance to copying the behavior of others: a review of public information use. *Behav Ecol Sociobiol.* 62:1–14.
- van Kesteren F, Delehanty B, Westrick SE, Palme R, Boonstra R, Lane JE, Boutin S, McAdam AG, Dantzer B. 2018. Experimental increases in glucocorticoids alter function of the HPA axis in wild red squirrels without negatively impacting survival and reproduction. *BioRxiv* [Preprint] April 26 2018 [cited February 13 2019]. doi:10.1101/309278.
- Wilson DR, Evans CS. 2012. Fowl communicate the size, speed and proximity of avian stimuli through graded structure in referential alarm calls. *Anim Behav.* 83:535–544.
- Wilson DR, Goble AR, Boutin S, Humphries MM, Coltman DW, Gorrell JC, Shonfield J, McAdam AG. 2015. Red squirrels use territorial vocalizations for kin discrimination. *Anim Behav.* 107:79–85.
- Yosida S, Okanoya K. 2009. Naked mole-rat is sensitive to social hierarchy encoded in antiphonal vocalization. *Ethol.* 113:113–131.
- Zuberbuhler K. 2009. Survivor signals: the biology and psychology of animal alarm calling. *Adv Study Behav.* 40:277–322.