Song overlapping, ambient noise and territorial aggression in great tits

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Abstract

Animals often communicate with each other in noisy environments where interference from the ambient noise and other signallers may reduce the effectiveness of signals. Signallers also may evolve behaviours to interfere with signals of their opponents, e.g. by temporally overlapping them with their own, such as the song overlapping behaviour that is seen in some songbirds during aggressive interactions. Song overlapping has been proposed to be a signal of aggressive intent although direct evidence for this hypothesis has been lacking. In the present paper we examined the question of whether song overlapping is correlated with aggressive behaviours in a population of great tits (Parus major) living in an urban-rural gradient. We also examined whether aggressive behaviours are correlated with the ambient noise levels. We found that overlapping was associated negatively while ambient noise levels were associated positively with aggressive behaviours males displayed against a simulated intruder. Ambient noise levels did not correlate with song rate, song duration or song overlapping. These results suggest that song overlapping of male great tits is not a signal of aggression and great tits in noisy urban habitats may display higher levels of aggression due to either interference of noise in aggressive communication or another indirect effect of noise.

Keywords: song overlapping, urbanization, noise, aggressive signalling, great tits
Introduction

The environment through which signals are sent and received is open to interference by either sources of noise or other signals traveling through the medium (Bee and Micheyl, 2008; Brumm, 2006; Gil and Brumm, 2014; Wiley, 2006). Signals should therefore evolve to reduce the level of interference from noise and other signals present in the environment (Endler, 1992; Endler and Basolo, 1998; Ey and Fischer, 2009). At the same time, signallers should evolve behaviours that would reduce the interference from noise or other signals, such as adjusting the timing of signalling to minimize overlapping with other signals (Brumm, 2006; Ficken et al., 1974; Gochfeld, 1978; Wasserman, 1977) or acoustic noise (Gil et al., 2014), adjusting the distance to the receivers (Halfwerk et al., 2012).

Generally speaking, signallers tend to space their signals temporally to minimize interference from other signalling individuals (Brumm, 2006; Ficken et al., 1974; Wasserman, 1977; Wilson et al., 2016). In some circumstances, however, actively overlapping the signal of another individual by starting to signal before the other signal has ended, may be a deliberate behaviour used as a signal itself. Indeed, overlapping has been proposed to be a signal of aggressive intent in songbirds (Dabelsteen et al., 1997; Mennill and Ratcliffe, 2004; Naguib and Mennill, 2010). Signals of aggressive intent (also called threat signals) are signals that are not costly to produce but carry information about the likelihood of escalation by the signaller (Caryl, 1979; Searcy et al., 2006). Under this hypothesis, overlapping and therefore interfering with the transmission of an opponent’s song may carry information to the opponent that the overlapping bird is ready to escalate an aggressive interaction.

Although song overlapping has been assumed to be an aggressive signal by a large number of studies (Naguib and Mennill, 2010), a review by Searcy and Beecher (2009, 2011) questioned whether song overlapping was a signal at all. Searcy and Beecher (2009) argued that a signalling behaviour should satisfy three empirical criteria to be considered a signal of aggressive intent: 1) the behaviour should occur more often in agonistic interactions than other non-agonistic interactions (context criterion); 2) the behaviour should correlate with other aggressive behaviours or predict a subsequent escalation (predictive criterion) and 3) the behaviour should elicit a differential reaction (either stronger or weaker) in receivers (response criterion).
Most studies of song overlapping have focused on the perspective of receivers, asking whether birds respond differentially when their songs being overlapped experimentally. These studies generally find changes in singing behaviours such as song rate and duration that are consistent with the singers attempting to avoid the interference due to being overlapped. Changes in aggressive behaviours such as attack or approach due to being overlapped however, are less consistent across studies (Naguib and Mennill, 2010; Searcy and Beecher, 2009).

Very few studies addressed the question whether song overlapping is correlated with aggressive behaviours or subsequent escalation, i.e. the predictive criterion (Baker et al., 2012; Fitzsimmons et al., 2008; van Dongen, 2006; Vehrencamp et al., 2007). Only one of these studies found potential support for overlapping being an aggressive signal: golden whistler males (*Pachycephala pectoralis*) were closer to the speaker when they overlapped the speaker than when they did not (van Dongen, 2006). Two other studies in black-capped chickadees (*Poecile atricapillus*) however, found no correlation between overlapping and concurrent aggressive behaviours or subsequent attack (Baker et al., 2012; Fitzsimmons et al., 2008). Finally a study in banded wrens (*Thryophilus pleurostictus*) found a negative correlation such that overlapping was correlated with lower levels of aggression (Vehrencamp et al., 2007).

In the present study we aim to assess the predictive criterion for song overlapping by examining the relationship between song overlapping and aggressive behaviours in great tits (*Parus major*). Although song overlapping has long been assumed to be an aggressive signal in great tits (Dabelsteen et al., 1996; Langemann et al., 2000; Peake et al., 2001), no previous study has asked whether this behaviour is correlated with the overlapping individual’s aggressive behaviours in this species.

A second aim of our study is to ask whether interference due to ambient noise also has an effect on aggressive behaviours. Although noise is a feature of natural habitats as well, the impact of noise in animal communication has garnered special attention due to the increasing levels of anthropogenic noise that affects social behaviour of wildlife (Gil and Brumm, 2014; Johnson and Munshi-South, 2017; Shannon et al., 2016). Anthropogenic noise may make signals less detectable (Kleist et al., 2016; Templeton et al., 2016) or less effective (Halfwerk et al., 2011). In response, animals may change the frequency, amplitude or length characteristics of their signals (Brumm and Todt, 2002; Slabbekoorn and den Boer-Visser,
2006; Wood and Yezerinac, 2006) or change their signalling or social behaviours (Halfwerk et al., 2012; Halfwerk and Slabbekoorn, 2009).

If noise interferes with effective signalling during aggressive interactions than we may expect signallers to display higher levels of aggression in noisier habitats. Consistent with this, in several species of songbirds birds living in noisier urban habitats display higher levels of aggression than birds living in rural habitats (Davies and Sewall, 2016; Evans et al., 2010; Fokidis et al., 2011; Foltz et al., 2015; Hardman and Dalesman, 2018). Multiple recent studies found more direct evidence for this hypothesis. In the first study male white-crowned sparrows (Zonotrichia leucophrys) holding territories with greater ambient noise displayed higher levels of aggression in response to song playback (Phillips and Derryberry, 2018). In another study, male house wrens (Troglodytes aedon) responded to male song playback near the nests with higher levels of aggression (Grabarczyk and Gill, 2019). Another recent study in chiffchaffs (Phylloscopus collybita) found that birds that were exposed to airport noise also responded more aggressively to conspecific playbacks compared to birds that were not exposed to noise. In contrast to these studies, a study examining territorial response in chipping sparrows (Spizella passerina) and spotted towhees (Pipilo maculatus) found that ambient noise was negatively correlated with approach latency to speaker in both of these species (Kleist et al., 2016). This effect may come about if ambient noise makes playbacks less detectable or harder to locate.

Noise also has an effect on singing behaviours, most notably in change of amplitude and frequency (Brumm and Slabbekoorn, 2005; Brumm and Zollinger, 2013), but also song rate (Brumm and Slater, 2006) and song duration (Ríos-Chelén et al., 2013). These changes may be adaptive in terms of ensuring the transmission of the signal to the receivers. Song overlapping may also change as a result of ambient noise: individuals may be less likely to overlap their opponent's songs in noisy environments if ambient noise already interferes with the transmission of the signals. To the best of our knowledge, no previous study examined whether song overlapping varied with ambient noise levels.

In the present study, we ask whether song overlapping and ambient noise levels are associated with territorial aggression in a population of great tits that live in habitats that range from urban to forest. As noted above, overlapping has been proposed to be an aggressive signal but evidence for the predictive criterion has been lacking. Previous research on this species showed that male great tits respond to anthropogenic noise by changing their song and behaviours: for instance, great tits in noisy urban areas sing songs with higher
minimum frequencies than great tits in quieter rural areas, which may benefit them as low-frequency songs are more prone to interference by anthropogenic noise (Hamao et al., 2011; Mockford and Marshall, 2009; Slabbekoorn and den Boer-Visser, 2006; Slabbekoorn and Peet, 2003). Urban-living great tits have also been found to be more aggressive than rural-living great tits (Hardman and Dalesman, 2018).

We studied great tits in an urban-forest gradient in Ankara, Turkey. In our population, located on the campus of Middle East Technical University, great tits hold territories throughout built-up areas as well as the forested areas adjoining the populated areas. Background noise in these areas vary significantly depending on proximity to buildings and roads, allowing us to ask whether ambient noise is correlated with aggression. We asked three specific questions: 1) Is song overlapping positively correlated with aggression across males? 2) Does ambient noise on territory predict aggressive behaviours? 3) Is ambient noise correlated with singing behaviours such as song overlapping, song rate, and song duration?

Methods

Study site and subjects

We studied great tits holding territories on the campus of Middle East Technical University, in suburban Ankara, Turkey (39°53'32"N, 32°47'03"E). Although the campus is located in a steppe habitat, it includes a large area that has been afforested mostly with conifers in the last six decades. Great tits nest in human structures on campus (e.g. access ports left open in telephone and electricity poles or cavities on the side of buildings) as well as nest boxes provided in the forested parts and natural cavities spread throughout the campus grounds. We located 42 territories of great tits by observation of singing posts or locating nests in the boxes or human structures. The birds were not captured at any time and therefore were not banded. The trials were carried out between 7 and 13 April 2019 at the start of the nesting period in the morning hours between 0600 and 1130. We avoided testing neighbouring males on the same day and carried out consecutive trials in locations that are outside of earshot of each other.

Playback Stimuli

We recorded male songs from great tits on campus in Spring 2018 and 2019 using a Marantz PMD660 or 661 recorder and a Sennheiser ME66/K6 shotgun microphone. Using the software Syrinx (John Burt, Seattle, WA) we viewed and selected high quality recordings of songs to create playback stimuli. After manually filtering out low-frequency noise (<~1000
Hz), we added a silent period to create a 7-second wave file. The average (±SD) playback
song duration was 2.99 (±0.42) seconds (range= 2.01-3.55 seconds). We created 24 playback
stimuli. For each subject we used a stimulus song that was recorded at least 1km away from
the subject’s territory. This constraint meant we used some stimulus songs multiple times for
different males (11 stimuli were used in two trials, 2 stimuli were used in three trials, and one
stimulus was used in 4 trials- the rest of the songs were used once).

Procedure

We placed a wireless speaker (Anker SoundCore, Anker, Inc) was placed face-up at a natural
perch inside the territory of the male (either 5 m from the nest if the nest was known or at a
location that was central to the various singing posts the male was observed to be singing).
We then set up flagging at 1m, 3m and 5m from the speaker on either side of the speaker to
help with distance estimation during the trial. The observers then stepped back to about 15 m
from the speaker. The playbacks were controlled from a smartphone connected to the speaker
via Bluetooth.

The trials were recorded with the same equipment as above. We started the trials by
playing back the stimulus song until the subject responded either by singing a song or
approaching the playback. After this first response, we continued the playback for another 3
minutes and we narrated the behaviour of the subject by noting each flight, distance to the
speaker and singing behaviours. The behaviours during this 3-minute period are the main
response variables of the study.

Noise measurements

After the trial, we removed the flagging, took a GPS reading (Garmin eTrek, Garmin Inc.) of
the trial location and carried out the ambient noise measurements using the method described
by Brumm (2004). Briefly, we took two measurements in each cardinal direction at the
location of the playback using a sound level meter (VLIKE VL6708, VLIKE Inc.) with A
weighting and fast response (125 msec) settings. In a subset of territories (n=32 out of 42) we
took a second measurement of ambient noise later in the day to assess how consistent
variation in the ambient noise was. For logistical reasons we could not take second noise
measurements in 10 territories. The first and second noise measurements were highly
correlated with each other (linear regression: β=0.94, SE= 0.17, t=5.46, p= 6.4 x 10^-6 ). We
used the first noise measurement in the analyses as this measurement was available for all
territories.
**Response measures**

Viewing the trial recordings in Syrinx, we extracted the following behaviours: flights, distance to speaker with each flight, and songs. Because the trial durations slightly varied across trials (ranging between 179 seconds to 192 seconds), we converted the counts of flights and songs into rates by dividing it with the trial duration. We also calculated the proportion of the trial subjects spent within 1 m of the speaker and noted the closest approach of the distance. For each song the subject sang, we determined whether the song overlapped with the playback song and calculated the proportion of the songs that the subject sang that overlapped the playback song. This measure therefore corrects for variation in song rate, but it is undefined for subjects that did not sing any songs (n=7). Because of variation in duty cycle (duration of stimulus with song/entire duration of stimulus) between stimulus songs, we also classified subjects based on whether the observed song overlapping (as defined above) was higher (n=11) or lower (n=24) than the duty cycle. We classified the subjects who did not sing any songs as a third category in this variable. Finally, for each subject we measured duration of each of their song and calculated the average song duration for each subject.

**Data Analysis**

The rates of flights, closest approach distance and proportion of time spent within 1m of speaker were all highly correlated with each other (Table 1). We therefore used a principle component analysis (PCA) to arrive at a single measure of aggressive response. The first component of the PCA (PCA1) explained 68% of variation and was taken as the aggressive score. The first component was positively correlated with flight rates and time spent within 1m, and negatively correlated with closest approach distance (Table 1).

We first ran two linear mixed models (LMM) with aggression scores (PCA1) as the response variable and stimulus song as a random factor. In the first model we entered the ambient noise levels measured after the trial as the fixed factor while in the second model we entered song overlapping as the fixed factor. We ran two separate models because song overlapping (as a proportion of subject songs that overlapped) was not defined for 7 subjects who did not sing any songs during the trial, thus reducing the sample size.

We then ran a parallel LMM to the first set of models with aggression scores again as response variable and stimulus song as random factor. In this model we entered ambient noise levels and the categorical overlapping variable as the predictor. This variable had three categories high-overlapping (overlapping rates higher than the stimulus duty cycle), low-
overlapping (overlapping rates lower than the stimulus duty cycle) or no song. This model included all subjects.

We additionally asked whether ambient noise was related to singing behaviours. To answer that question we ran LMMs on song rate, song overlapping and average song duration as response variables, and ambient noise levels as predictor variables, with stimulus song as a random factor, to determine whether ambient noise had an effect on other singing behaviours. Finally, we compared the observed level of song overlapping with the expected levels (represented by the duty cycle), by carrying out a paired t-test. The analyses were carried out in R using the package nlme (Pinheiro et al., 2017; R Core Team, 2012).

Results

Contrary to the hypothesis that song overlapping is a signal of aggressive intent, overlapping (proportion of subject songs that overlapped the playback songs) was negatively related to the aggression scores: high overlapping was indicative of lower aggression scores (Table 2, Figure 1). Ambient noise was positively related to aggression scores: birds in noisier territories were more aggressive in response to the playback (Table 2, Figure 2). In the model with the categorical overlapping variable (Table 3), both overlapping and ambient noise significantly predicted attack: birds that overlapped the playback less than expected were more aggressive compared to high-overlapping birds (unpaired t-test: t(33)=2.78, p=0.0088). The most aggressive birds were birds who did not sing any songs who were more aggressive than low-overlapping birds (unpaired t-test: t(29)=2.10, p=0.044; Figure 3).

Ambient noise was not related to song overlapping, song rate or average song duration (Table 4). The observed levels of overlapping (mean ± SD: 0.33 ± 0.29) was lower than the duty cycle of stimulus songs (0.42 ± 0.06) although not significantly so; unpaired t-test unequal variances: t(34)=-1.77, p=0.085.

Discussion

In this study we asked whether song overlapping and ambient noise was related to territorial aggression in male great tits in a population that spans an urban-rural gradient. We found that song overlapping was negatively related to aggressive behaviours suggesting that high levels of overlapping is not a signal males use to threaten opponents. We also found that ambient noise levels were positively related to aggression scores, providing support for the hypothesis that high ambient noise levels may select for higher aggression levels. Nonetheless, ambient
noise levels were not related to singing behaviours and parameters such as song rate, duration or overlapping.

*Is song overlapping an aggressive signal?*

The present results suggest that song overlapping is not necessarily a reliable signal of aggressive intent in great tits. Our finding of a negative correlation between aggression and song overlapping is similar to what Vehrencamp and colleagues (2007) found in banded wrens. In another species, the black-capped chickadees, song overlapping was not associated with aggressive behaviours or attack (Baker et al., 2012). Taken together, these results imply that song overlapping does not necessarily carry information regarding high likelihood of escalation to the receivers although it remains to be seen whether the negative conclusion will apply generally to other species. Additionally, song overlapping tended to happen slightly lower than expected based on the duty cycle of stimulus songs but this difference was not significant, indicating that overlapping may not be a deliberate behaviour in any case.

It is worth noting that there is some debate recently about what the correct null expectation should be in studies of song overlapping (Masco et al., 2016; Naguib and Mennill, 2010; Searcy and Beecher, 2009, 2011). We used stimulus duty cycle (the ratio of the stimulus song duration over the sum of the stimulus song duration and silent period between two stimulus songs) as the expected levels simply because it does not rely on any additional assumptions. Choosing a different “null” model, with varying assumptions, may yield a result indicating a significant difference between expected and observed levels of overlapping (Masco et al., 2016). Nonetheless, whatever the null model, if song overlapping is a deliberate behaviour that individuals use to signal their intent to escalate, individual variation in song overlapping should correlate positively with aggressive behaviours.

If song overlapping is not an aggressive signal in great tits, what does it indicate? One possible answer is that high levels of overlapping may simply indicate that the overlapping bird is not engaged with the intruder, whereas low levels of overlapping may indicate a closer engagement. In fact, birds who approached the speaker more closely and flew around the speaker more in search of the opponent may be avoiding overlapping the opponent as an assessment strategy (Arnott and Elwood, 2009). Great tit song has been found to contain information regarding the singer’s traits such as age and condition (Rivera-Gutierrez et al., 2010). Thus, it may be advantageous for receivers that intend to engage the intruder at close range to listen carefully to the song features of the opponent. This listening strategy may
explain the low rates of overlapping among aggressive individuals as well as the fact that birds that do not sing are the most aggressive group.

It is also possible that non-overlapping the intruder may simply be due to a search strategy to locate the opponent acoustically while the opponent is singing. Under this hypothesis, the males are avoiding overlapping the intruder because they are trying to figure out where the intruder is. Indeed, singing while the opponent is also singing is likely to interfere with the processing of spatial information that can be obtained from the song of the intruder. If that is the case, then birds that are aggressively responding to their opponents and therefore more motivated to locate them would be less likely to overlap their opponent’s songs. This hypothesis is essentially an extension of the “readiness hypothesis” that was proposed for explaining the occurrence of low amplitude “soft” songs that is used as a signal of aggressive intent in several songbirds (Akçay and Beecher, 2012; Akçay et al., 2011). The readiness hypothesis posits that for soft songs, the low amplitude is simply a by-product of the fact that the beak and body movements required to sing loudly might interfere with visual tracking of an opponent. Therefore, in an aggressive interaction birds that intend to escalate might be signing with their beak closed, which leads to lower amplitudes (Akçay and Beecher, 2012; Goller et al., 2004). Similarly, singing while the intruder is also singing (i.e. overlapping) may interfere with localizing of the opponent acoustically and birds that intend to escalate the interaction may therefore avoid overlapping the opponent’s song. This hypothesis can be tested with an experiment in which the simulated intruder can be either located by the subject either easily or with difficulty.

Another possibility is that overlapping, instead of being a signal of aggressive intent, may be correlated with more permanent traits of the signallers, such as some aspect of genetic quality (Helfer and Osiejuk, 2015). In their recent review, Helfer and Osiejuk (2015) presented several hypotheses to that positing that song overlapping is a signal of male quality due to some a physical or physiological constraint or a handicapping mechanism. However, given that song overlapping is a rather flexible behaviour that primarily depends on the chosen timing of the singing it seems unlikely that it would be subject to a physical or physiological constraint or handicapping mechanism. Interestingly, a study on great tits found that great tits that were exposed to an ectoparasite as nestlings were found to be less likely to overlap the song playback carried out in the territory after they were recruited to the population (Bischoff et al., 2009). Song overlapping however, was not correlated with the condition of the male as a nestling, and there was only a trend towards a positive association.
between overlap and male condition as an adult. Thus, although these data suggest a potential
developmental constraint on song overlapping, the mechanism is unclear. Bischoff et al.
(2009) do not present any information on aggressive behaviours such as approach, thus it is
not clear whether the developmental handicap was associated with aggressive behaviours as
well. In summary, while it remains possible that overlapping reflects either male quality,
condition or developmental history, more data are clearly needed to test these possibilities.

Noise and aggressive behaviours

Ambient noise was positively correlated with aggressive behaviours in the present study,
consistent with the hypothesis that noise may play a significant role in determining
aggressive phenotypes observed in urban populations of birds (Davies and Sewall, 2016;
Evans et al., 2010; Hardman and Dalesman, 2018). The effect of noise on aggression may
come about via multiple mechanisms. First, noise may disrupt effective communication
(McMullen et al., 2014; Ríos-Chelén et al., 2015). When effective communication is
disrupted, aggressive behaviours may increase in prevalence (Kareklas et al., 2019; Logue et
al., 2010). Interestingly, unlike other species (Brumm and Slater, 2006; Ríos-Chelén et al.,
2013) great tits in our population did not show any evidence that they change their song rate,
average song duration or overlapping depending on ambient noise levels. Given that previous
research showed that songs in great tits and other songbirds become less effective in urban
noise (Halfwerk et al., 2011; Luther et al., 2015), this finding suggests communication may
indeed be hampered in noisier territories.

A second, non-mutually exclusive possibility is that noise may lead to higher levels of
physiological arousal or stress which may increase aggression, as has been documented in
humans (Baron and Richardson, 2004). In particular, exposure to noise may elevate baseline
levels of glucocorticoids (Creel et al., 2002; Crino et al., 2011; Davies et al., 2017; but see
Kleist et al., 2018) which in turn may lead to a higher level of aggression in individuals living
in noisier areas. Interestingly, in a recent study on urban and rural song sparrows (Melospiza
melodia), Davies and colleagues (2018) found that baseline levels of corticosterone (the main
avian glucocorticoid) in blood samples after a simulated territorial intrusion did not differ
between urban and rural birds, and the levels of corticosterone were positively related to
territorial aggression males displayed. For a given level of plasma corticosterone however,
urban birds displayed more aggression than rural birds, suggesting that urban birds
potentially show differences in glucocorticoid signalling from rural birds. Thus, the exact
physiological mechanisms that determines the relationship between urbanization and noise on
the one hand, and aggression on the other is yet unclear (Bonier, 2012). Further experimental studies are needed to quantify the effect of noise on plasticity at behavioural and physiological levels to tests hypotheses on the physiological mechanisms of aggression under noisy conditions.

Conclusion

In summary our findings on song overlapping suggest that this signal is unlikely to be a signal of aggressive intent, and researchers should pursue new hypotheses for this interesting behaviour. Our findings on ambient noise meanwhile, corroborate the recent studies linking noise and aggression and suggest new avenues to investigate the effect of anthropogenic noise on animal social behaviour and the physiological mechanisms underlying it. Together, the current study highlights how animals may manage interference from noise or other signallers in their social interactions.
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Figures.

Figure 1. Scatterplot of song overlapping (proportion of the subject’s songs that started while the playback song was playing), and aggression scores (PCA 1).
Figure 2. Scatterplot of ambient noise levels (average of 8 measurements with A settings, dB) and aggression scores (PCA1)
Figure 3. Aggression scores of subjects a) overlapping the playback songs at more than the duty cycle (n=11), less than the duty cycle (n=24), singing no song (n=7).
Table 1: Pearson correlation coefficients (p-values) for the correlation between the aggressive response variables and their loadings to the first component of principle component analysis (PCA).

|                                           | Flight rate | Proportion of trial spent within 1m | Closest approach |
|-------------------------------------------|-------------|-------------------------------------|------------------|
| Flight rate                               | 0.60 (2x10^-5) | -0.57 (7x10^-5)                     |                  |
| Proportion of trial spent within 1m       | 0.42 (0.005)  | -0.551                              |                  |
| Loading Coefficient                       | 0.614        | 0.564                               | -0.551           |

Table 2. Linear mixed models (LMM) with stimulus song as random factor and aggression scores as response variable. The first model has song overlapping as the fixed factor (n=35 subjects, 7 subjects are excluded because they did not sing a song, and proportion of their songs that overlapped the stimulus song was therefore undefined). The second model has ambient noise as the fixed factor (n=42 subjects).

| Model                  | Coefficient (SE) | $\chi^2$ | p-value |
|------------------------|------------------|----------|---------|
| Song overlapping       | -1.89 (0.76)     | 6.27     | 0.012   |
| Ambient noise          | 0.09 (0.035)     | 6.94     | 0.0083  |

Table 3. Linear mixed model on aggression scores with the categorical overlap variable and ambient noise as predictor variables. Stimulus song is a random factor.

|                         | $\chi^2$ | p-value   |
|-------------------------|----------|-----------|
| Overlap                 | 16.41    | 0.00027   |
| Ambient noise           | 6.01     | 0.0083    |

Table 4. Linear mixed models on song rate, average song duration and song overlapping as response variables and ambient noise as predictor variable. Stimulus song is a random factor in all models.

| Model                  | $\chi^2$ | p-value |
|------------------------|----------|---------|
| Song rate              | 0.063    | 0.80    |
| Song duration          | 0.69     | 0.79    |
| Song overlapping       | 2.18     | 0.13    |