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The flexible listener: exploring zebra finch sensitivity to spectral and temporal sound features

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The Role of Spectral Features and Song Duration in Zebra Finch Song Recognition

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ABSTRACT

Zebra finch song perception is assumed to primarily involve a high sensitivity to fine spectral features of song elements while other features like element sequence and song duration do not seem to have a notable effect. However, the specific features that zebra finches focus on when identifying or discriminating sounds may not be as fixed as seems to be assumed and might depend on the characteristics of the stimuli. This apparent flexibility in auditory processing, along with the potential salience of differences in song duration for song perception, highlights the need for systematic research on the acoustic parameters that zebra finches can use to differentiate between songs. By employing a Go-Left/Go-Right operant task, we examined whether and how differences in song duration affect zebra finches' relative sensitivity for spectral features and duration in song recognition. Two groups of zebra finches were trained in a Go-Left/Go-Right operant task to discriminate either between two songs with similar durations ("Equal-duration group") or two songs with different durations ("Unequal-duration group"). We assessed to what extent the birds in the two experimental groups attend to the spectral characteristics and the absolute duration of the songs by measuring the responses to test stimuli consisting of spectral modifications or temporal changes. Our results showed that zebra finches use both spectral features and song duration to discriminate between two songs, but the importance of these acoustic parameters depended on whether the songs differed in duration or not. When duration can be used as an additional feature to distinguish two songs, spectral features have a less prominent role. This outcome shows that zebra finches have cognitive flexibility in their attention to different acoustic parameters.

INTRODUCTION

Birdsongs convey important information that varies from individual identity to information about sex, age, individual quality, or motivation. Meaningful communication requires that receivers be able to perceive and process the acoustic variation in songs. On the one hand, regardless of external conditions that may affect the transmission of song features, a receiver has to recognize a song as coming from the same singer. At the same time, the receiver must be able to discern meaningful variations within songs produced by the same singer, as well as being able to distinguish between songs from different individuals. This raises the question of the cognitive mechanisms through which songbirds recognize and classify songs, and discriminate between different songs and song variants. Experimental studies have addressed this topic in various ways, ranging from field studies using playback to psychophysical laboratory experiments using operant discrimination paradigms. Field studies examined, for example, the characteristics birds employ to recognize conspecific songs or to discriminate between conspecific and heterospecific ones (e.g., Nelson, 1989; Dabelsteen & Pedersen, 1992; Naugler & Ratcliffe, 1992). Psychophysical studies have been used to investigate the hearing ranges and the abilities of birds to detect specific details in the spectral or temporal structure of songs (e.g., Kreutzer et al., 1990; MacDougall-Shackleton & Hulse, 1996; Tu & Dooling, 2012; Neilans et al., 2010; Dooling & Prior, 2017). Such studies have provided important insights in the mechanisms underlying auditory perception and communication in birds. At the same time, studies on avian sound perception are relevant from a comparative perspective as they can reveal the presence of both similarities and differences in the acoustic features that are salient or noticeable by humans and those to which birds attend (e.g., Hulse et al., 1984; Bregman et al., 2012; Hoeschele, 2017; Dooling & Prior, 2017).

Over the years, the zebra finch (*Taeniopygia guttata*) has emerged as a model species for examining the processing of complexly structured auditory stimuli at the level of behaviour as well as its underlying neurobiology. One area of research concerns the features that zebra finches can or do use to recognize or discriminate between songs. These features are often examined by using operant discrimination tasks. For instance, using a design in which zebra finches were first trained to respond to a single song type and not to respond to deviations, Dooling and collaborators (Dooling et al., 2002; Lohr et al., 2006; Vernaleo et al., 2010; Vernaleo & Dooling, 2011; Lawson et al., 2018; Prior et al., 2018a; Prior et al., 2018b; Fishbein et al. 2021) examined the salience of various types of song changes on the

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identification of the target song. They showed that zebra finches are quite sensitive to changes in the spectro-temporal structure of syllables but relatively insensitive to changes in syllable order in zebra finch song motifs. From these studies, they concluded that zebra finches primarily attend to spectral details such as the temporal fine structure (phase in the waveform over extremely short periods) within individual syllables. Also, several other studies (e.g., Uno et al., 1997; Vignal & Mathevon, 2011; Geberzahn & Derégnaucourt, 2020; Mol et al., 2021) indicated the prominent importance of spectral features for vocal discrimination in zebra finches, with low-frequency harmonics more important for song identification than high-frequency ones (Dent et al., 2016). A prominence of spectral features over syllable sequence for discriminating songs was also shown by Braaten et al. (2006) using a Go/Nogo paradigm. In another study, Nagel et al. (2010) trained adult female zebra finches to perform a classification task in a two-alternative forced choice paradigm to investigate the role of three acoustic parameters (pitch, tempo, and amplitude) in discriminating between two male songs. Small changes in pitch ($\pm 2\%$) already affected song discrimination, while tempo alterations affected song discrimination only when these were substantial ($> 32\%$).

The above studies suggest that the main factors involved in sound perception in zebra finches are known and predictable: a high sensitivity for fine spectral features of acoustic stimuli with substantially less, if any, impact on other parameters, such as tempo (speed) or song duration. However, several other findings suggest that the features to which zebra finches attend when identifying or discriminating songs or other auditory stimuli are not as fixed as the experiments mentioned above suggest and may depend on the characteristics of the stimuli. For instance, in contrast to the study by Nagel et al. (2010), which suggested that zebra finches hardly attend to tempo changes of auditory stimuli equal to or less than 32%, other experiments demonstrated that zebra finches do respond to small tempo changes when two series of identical sound pulses could only be differentiated by attending to temporal features (van der Aa et al., 2015; ten Cate et al., 2016). Here a 25% change in tempo substantially reduced stimulus discrimination. The contrast of this finding with the limited impact of any tempo changes on song identification, as obtained by Nagel et al. (2010), may arise because Nagel et al. (2010) used songs with a similar song duration. If the duration of songs is similar, then duration might be an irrelevant parameter for song identification and therefore ignored. Spectral features are then the main distinguishing parameter, and zebra finches might focus their attention on such features to identify songs. This might explain a limited effect of tempo changes on song identification compared to the discrimination of auditory stimuli consisting

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of identical elements, differing in tempo only. Similarly, syllable sequence is not a prominent parameter when zebra finches are trained to discriminate two syllable strings consisting of different song syllables. However, when the two strings consist of the same syllables but in a different sequence, zebra finches attend to the sequence in addition to the spectral structure of the syllables (Ning et al., 2023). This indicates that zebra finches are flexible in the auditory parameters they attend to and use those acoustic features that allow them to differentiate between the stimuli. This was also suggested by a study in which zebra finches were trained to discriminate between two sets of artificial vowel-like harmonic elements (Burgering et al., 2019). For one group of birds, the distinguishing feature of the spectra was the fundamental frequency (pitch), while for the other group this was the relative energy distribution over the harmonic spectrum across the elements, indicated as the “spectral envelope”. Probe tests showed that the first group maintained the discrimination when the energy distribution over the spectra was changed, but the fundamental frequencies remained the same. The second group of birds ignored changes in the fundamental frequency of the spectra but maintained the discrimination when the harmonic sounds were replaced by a noise-vocoded sound. Such a manipulation divides the original sound into distinct frequency bands and replaces the spectral variation within each band by a noise signal with the same amplitude. The results of this experiment thus show that zebra finches can either ignore or use the fundamental frequency or the harmonic structure of the sound depending on which is relevant for acoustic discrimination. It also shows that zebra finches can attend to the shape of the spectral envelope, something that had not been tested before in zebra finches, but which had been demonstrated by Bregman et al. (2016) for starlings discriminating among more complex tone sequences. Bregman et al. (2016) suggest that the spectral envelope governs avian tone sequence recognition. The importance of this feature may long have gone unnoticed as many experiments on avian pitch perception used pure tones, for which the spectral band envelope corresponds directly to pitch. The findings of Burgering et al. (2019) and Bregman et al. (2016) may indicate that the sensitivity of zebra finches to pitch changes in songs need not necessarily indicate a sensitivity to pitch, but alternatively might result from being sensitive to the spectral envelope, something that has so far not been tested for song stimuli.

The apparent flexibility in the features used during auditory processing shown by zebra finches and the potential role of duration and spectral envelopes in song perception call for further research on the acoustic parameters that zebra finches can or do use to distinguish between songs. The present study aims at exploring these parameters. Two groups of zebra finches

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were trained to discriminate two songs in a Go-left/Go-right task. For one group these songs were equal in duration (Equal-duration group), for the other group they were unequal in duration (Unequal-duration group). After being trained, zebra finches were tested with modified versions of training songs that were changed in one of the following ways: 1) increasing or decreasing the tempo, thus affecting the duration; 2) raising or lowering the pitch; 3) moving the entire song up in the frequency spectrum; or 4) replacing the harmonic spectrum by a noise-vocoded version. This design allows us to examine two factors. The first one is whether song duration is used as an additional factor to spectral features when zebra finches are trained to discriminate two songs that differ in duration. If the duration is used as an additional factor, we expect that learning will be easier and hence the training phase will be shorter when learning to discriminate between songs of different compared to similar duration. We also expect that the relative impact of modifying spectral features versus temporal ones on the ability to recognize and discriminate among songs will differ depending on whether training songs differ in duration. For birds of the Equal-duration group, song duration is not a distinguishing factor between the training songs, while it is for birds of the Unequal-duration group. Therefore, we expect that zebra finches trained to discriminate two songs of equal duration will be less sensitive to tempo changes of the songs than zebra finches trained with songs of different duration. In contrast, we expect that birds from the Equal-duration group will be more sensitive to changes in the spectral domain, because they can only use spectral features to discriminate the training songs. Thus, the relative impact of tempo changes and spectral changes is expected to differ between the two experimental groups. The second factor we examine is the relevance of the spectral envelope *vs* pitch in song discrimination. If zebra finches, like starlings, attend to the spectral envelope rather than pitch for song recognition, we expect that vocoded songs may be easier to recognize than songs with pitch changes or songs moved up in frequency. In this case we have no reason to expect a difference between the Equal-duration and the Unequal-duration groups. However, for each group we expect that if the birds attend more to the spectral envelope than to pitch, the vocoded version of the song will be considered more similar to the training songs than songs in which the pitch or frequency profile has been shifted.

METHODS

Subjects

We tested a total of 28 zebra finches (14 males, and 14 females; ages 215-720 days post hatching) originating from the in-house breeding colony at Leiden University. Before the experiment, the birds lived in single-sex groups of about 15 to 30 individuals in aviaries (2m × 2m × 1.5m), in which food and water were available ad libitum. The birds were divided equally between two experimental groups, each consisting of seven males and seven females. Each group was trained with a different set of stimuli, and within each group half of the birds got one set of test stimuli ('series 1') and half another set of test stimuli ('series 2'), hence resulting in a total of four subgroups, each consisting of seven birds.

Operant conditioning cage

The birds were trained and tested individually in an operant conditioning cage (Skinner box) (70x30x45 cm) containing 3 pecking keys (sensors) with a red LED light at the top/bottom of each sensor (Fig. 1). Each operant cage was situated in a separate sound-attenuated chamber. The chamber was illuminated by a fluorescent lamp (Phillips Master TL-D 90 DeLuxe 18W/965, The Netherlands), which emitted a daylight spectrum following a 13.5-h/10.5-h light/dark schedule. Sound stimuli were played through a speaker (Vifa MG10SD09-08; frequency range 100 – 15000 Hz) 1 meter above the Skinner box. The volume of the speaker was adjusted to ensure that the sound amplitude in the Skinner box was approximately 65 dB (measured by an SPL meter - RION NL 15, RION), a level comparable to what the bird would be exposed to from a singing conspecific at the location of the bird. Sensors (S1, S2, S3), lamp, food hatch and speaker were connected to operant conditioning controller that also registered all sensor pecks.

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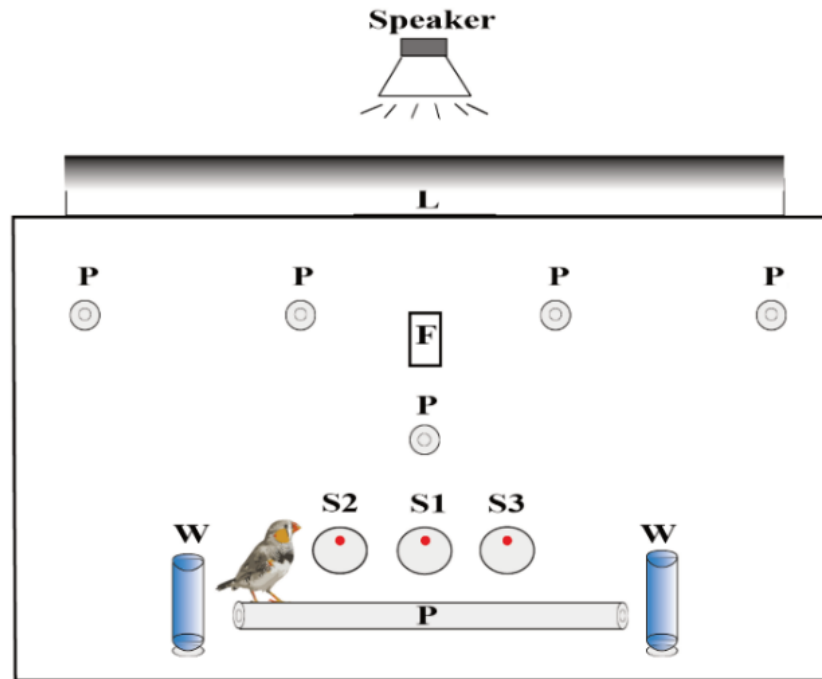


Figure 1. Schematic front view of the operant conditioning apparatus (Skinner box) used for the experiment. A speaker (top of figure) is suspended from the ceiling above the cage. Within the cage, there are several perches (P) for the bird to sit on, a food hatch (F) located in the upper middle of the back panel, a lamp (L) is placed at the top of the cage. Two tubes with ad libitum water (W) are placed symmetrically on two sides of the cage, and three sensors (S1, S2, S3) with red LEDs are lined horizontally in the lower middle of the back panel.

Stimuli

Training stimuli

A total of 24 natural song motifs were used. The song motifs were extracted from representative recordings of adult males from our breeding colony, but whose vocalizations had not been heard before by birds in this study. The training stimuli in this experiment were 14 stimulus pairs (seven pairs for each experimental group), each consisting of two different songs. Every stimulus pair was used twice, for two separate subgroups of birds ($N = 7$ birds/group). The two subgroups of birds per training stimulus pair were subjected to different series of test sounds – one subgroup to test series 1 and the other to test series 2 (see below). Of the 14 stimulus pairs, seven pairs consisted of songs of approximately equal duration, in which the shortest song always differed less than 5% from the duration of the longest song in a pair (mean duration of the shortest song was $98.21\% \pm 1.45\%$ of the duration of the longest song). The group trained with these stimuli will be indicated as the ‘Equal-duration group’. For the other seven pairs the songs were of unequal duration, with the duration of song A being

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approximately 1.5 times longer than its paired song B (mean: $148.43\% \pm 6.50\%$). The group trained with these songs will be indicated as the ‘Unequal-duration group’ (Fig. 2). Hence the experimental structure was that both the ‘Equal’ and the ‘Unequal’ group consisted of two subgroups of seven birds, each trained with the same stimulus set, but tested with a different set of test stimuli.

Within each song stimulus, the same motif was repeated three times with a silent gap between the motifs, thus simulating a natural song sequence. When played, the motifs were normalized such that the average intensity (RMS - calculated over the total duration of the stimulus) was the same for the two stimuli within a set but the amplitude variation of the original male zebra finch song was preserved. All training stimuli were bandpass-filtered between 380Hz and 22.5kHz. The two stimuli from each training stimulus set were visually selected to differ in the spectral structure of the syllables (Fig. 2). All training stimuli were cut, synthesized, and filtered using Praat (version 6.0.54). The amplitude of each stimulus was adjusted by using the “Normalize” feature in Audacity (version 2.3.0).

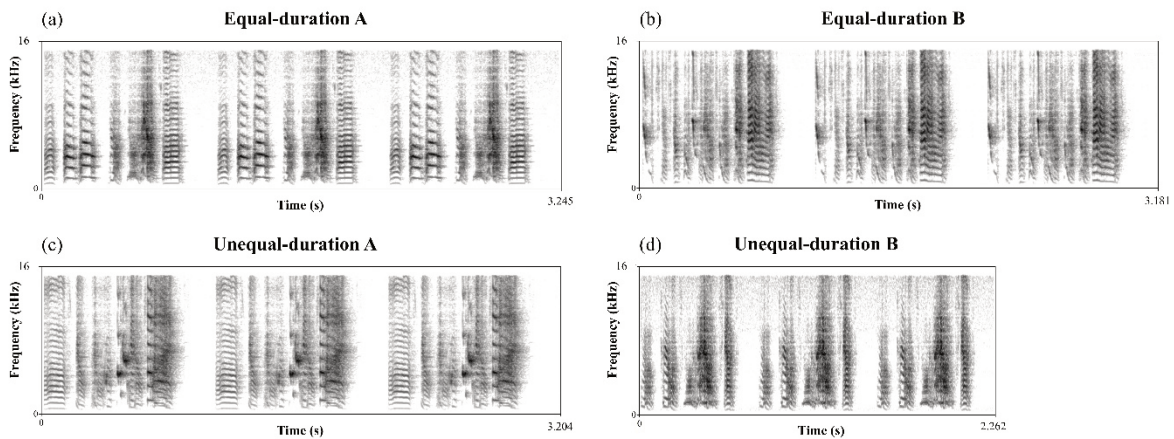


Figure 2. Spectrogram samples of training stimuli. Songs Equal-duration A (a) and Equal-duration B (b) form a pair of training stimuli used in the Equal-duration group, while songs Unequal-duration A (c) and Unequal-duration B (d) form a pair of training stimuli used in the Unequal-duration group.

Test stimuli

To test the impact of specific parameters that the birds may have used to discriminate the training stimuli, they were tested with modified versions of the training stimuli, which were grouped into two series of test stimuli (Table 1). The two series differed from each other in how strongly they modified specific parameters of the training stimuli. We expected that a stronger modification would have a stronger impact on song discrimination. Each subgroup of

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birds was tested with one series of sounds only. We used the Praat Vocal Toolkit (A Praat plugin with automated scripts for voice processing, www.praatvocaltoolkit.com) to edit each original training stimulus to produce a version with either spectral features or the tempo was changed. For both the Equal-duration and the Unequal-duration training group, the test stimuli were always modified from the training stimuli in an identical way. We used the following set of test stimuli (Fig. 3, Table 1):

Table 1 Overview of test stimuli used for the two experimental subgroups

	Frequency -shifted	Pitch-shifted upward	Pitch-shifted downward	Duration stretched	Duration compressed	Noise-vocoded
1st series	+1500Hz	+20%	-20%	+50%	-50%	Spectral envelope maintained but spectral contour averaged
2nd series	+500Hz	+8%	-8%	+20%	-20%	Spectral envelope & spectral contour maintained

Note: Two test series were used for the subgroups of both the Equal-duration and Unequal-duration experimental group of birds. The test stimuli differed in the degree to which pitch and duration were modified, which was more strongly changed in series 1 than in series 2 and in the scripts used for vocoding (series 1: vocoded version according to the script by Chris Darwin; series 2: vocoded version according to the script by Matt Winn. See text for details).

- Frequency-shifted – For this stimulus the whole frequency spectrum was shifted upwards linearly. By this manipulation, the harmonic relations between the frequencies are no longer preserved. This was obtained by using a Fresh plugin of Audacity (version 2.3.0 - full buckets frequency shifter, www.fullbucket.de/music), adding a fixed value to the frequency of each component of the original sound signal. For subgroup of birds tested with series 1 this value was 1500 Hz and for subgroup of birds tested with series 2 this was 500 Hz.

- Pitch-shifted – The frequency spectrum was stretched or compressed on a log scale to produce a version in which the harmonic relationship between the frequencies in the song remain the same, but their absolute frequencies were changed. This version of the target sound was synthesized using the “Change vocal trace” script of the Praat Vocal Toolkit by entering the specific formant shift ratio value in the options displayed in running this script. For the subgroup of birds tested with series 1, the frequency spectrum was stretched or compressed by 20%, and for the subgroup of birds tested with series 2 it was 8%. The choice of the values of 8% and 20% was based on the study by Nagel et al. (2010), in which an 8% change resulted in a reduced discrimination between two songs, although they were still discriminated above

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chance, while a 50% change resulted in lack of discrimination. The 20% value thus was intermediate between these.

- Time-scaled – The duration of the whole song was stretched or compressed proportionally without any change in the frequency domain. The “change duration” script of Praat Vocal Toolkit was applied to obtain stretched and compressed song versions. For the subgroup of birds tested with series 1, the duration was stretched or compressed by 50%, and for the subgroup of birds tested with series 2 it was 20%. Here also the values of 20% and 50% were chosen based on the study by Nagel et al. (2010) in which a 20% change did not affect the degree of song discrimination, while a 50% change reduced (but not eliminated) the discrimination.

- Noise-vocoded – This modification maintains the spectral envelope (the overall shape of the frequency spectrum) of the elements within the motif, but averages the energy within specific frequency bands, thus removing any harmonic structure. To construct these stimuli, we used two different scripts to synthesize a vocoded morph of training stimuli: for the subgroup of birds tested with series 1, we used the modified Chris Darwin vocoded script (for the original version, see http://www.lifesci.sussex.ac.uk/home/Chris_Darwin/Praatscripts/Shannon) which also removed the within-syllable spectral contour (the shape of the sound’s frequency components over time) of the song syllables (referred to as ‘Contour-averaged Vocoded’), and for the subgroup of birds tested with series 2 we used the Matt Winn’s Praat vocoded script (http://www.mattwinn.com/praat/vocode_all_selected_40.txt) which maintained the within-syllable spectral contour (referred to as ‘Contour-maintained Vocoded’). Both these two scripts were set to divide cut-off frequency bandwidths equally for 15 bands contiguous with smooth transitions (1000Hz bandwidth for one noise-vocoded band).

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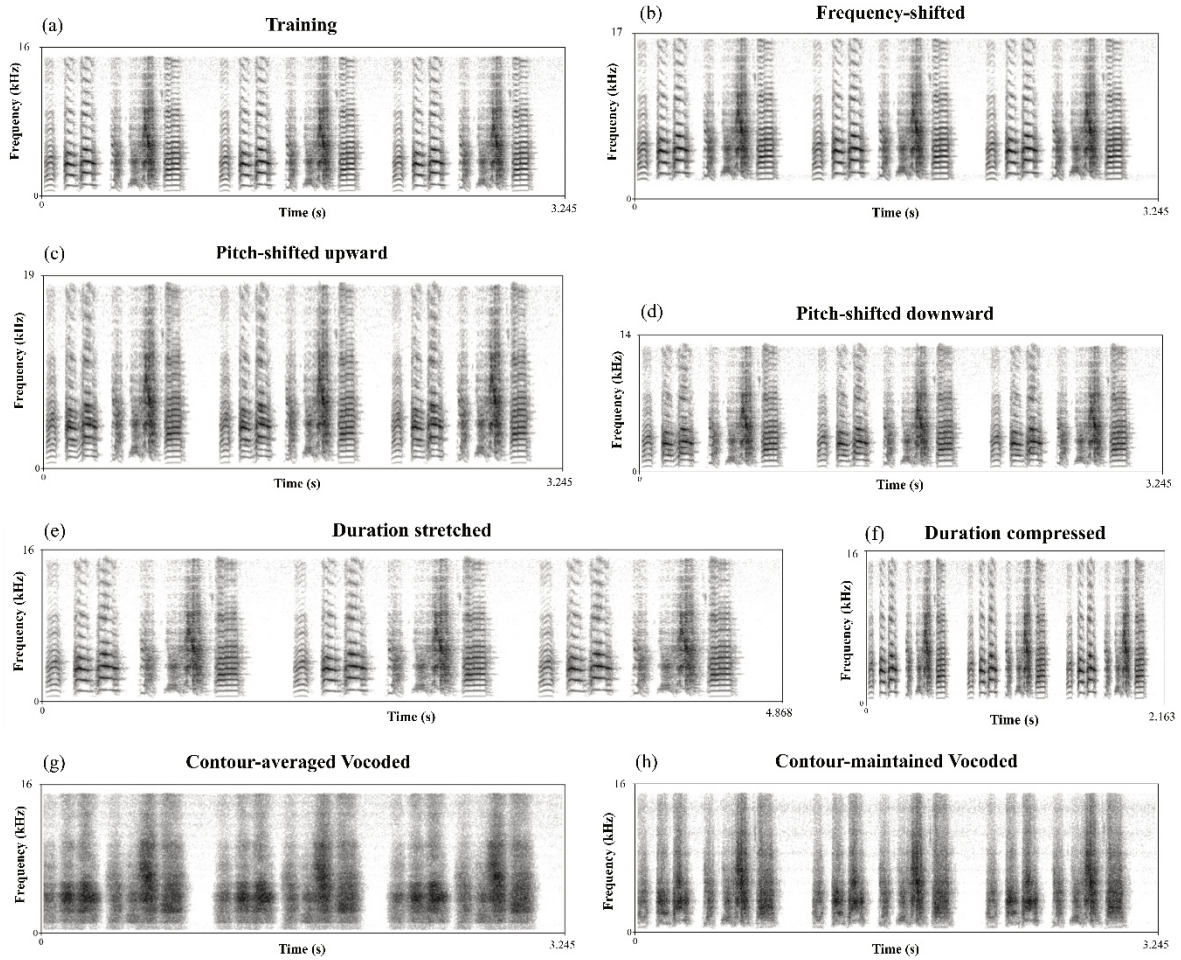


Figure 3. Examples of stimuli used in the test series, showing (a) the **Training stimulus**, and its modified versions. The whole frequency spectrum of (b) the **Frequency-shifted** version was shifted upwards by 1500 Hz. The frequency spectrum of the **Pitch-shifted** stimulus was either (c) stretched (+20%) or (d) compressed proportionally (-20%). The duration of the **Time-scaled** stimulus was either (e) stretched (+50%) or (f) compressed (-50%). The **Noise-vocoded** versions were produced by using two scripts – (g) the modified Chris Darwin vocoded script (Contour-averaged Vocoded) and (h) the Matt Winn’s Praat vocoded script (Contour-maintained Vocoded).

Procedure

We used a Go-left/Go-right paradigm for training and testing. The procedure consisted of five phases: acclimation, pre-training, discrimination training, transition, and probe testing. The birds stayed in the Skinner Boxes during all phases of the experiment.

Acclimation phase

In the acclimation phase the birds were moved to the Skinner boxes (See Fig. 1). The food hatch remained open, so food was freely accessible in a container behind the hatch. The LED

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lights on the sensors were on. The goal of this phase was to acclimate the bird to the cage and show where to find food. The bird might also already learn to peck the sensors spontaneously. If in this stage the central sensor, S1, was stimulated by pecking, it would play song A or song B with a 50% chance on each. The side sensor S2 produced one of the two songs, and the other side sensor S3 produced the other song. The red LEDs of all three sensors were illuminated to attract the attention of the bird. After several hours to one day, with a median value of 26 (IQR 18-28) hours, the next phase started by closing the food hatch.

Pre-training phase

The goal here was to familiarize and teach the bird training procedures. In this phase, the food hatch was closed, and the bird had to learn to peck all three sensors. Pecking the sensors in this phase led to the following effects: S1 (middle sensor) = sound A or B (no food), S2 (side sensor) = sound A + food hatch open (duration 12 seconds), S3 (side sensor) = Sound B + Food hatch open (12 seconds). This continued until the bird had learned to peck at each sensor, and that pecking the sensors resulted in access to the food. The bird might also already learn at this stage which song was related to S2 or S3. This process took several days, with a median value of 95 (IQR 68-122) hours. If the bird did not peck the sensor spontaneously, the experimenter could turn on/off the LED to make it flash to stimulate the bird to pay attention to the sensor. Once the bird started pecking all the sensors regularly (i.e., pecking each of the three keys over 50 times in one day) for a day, the discrimination training phase began.

Discrimination training

In this phase, the bird had to learn to peck the sensor in the middle to elicit the playback sound, followed by pecking the sensor on the left or right, depending on the playback sound. If the bird pecked the sensor linked to the particular stimulus being played, a response was rewarded with 12 sec access to food. If the wrong sensor was pecked, the light was off for three seconds. Before any sensor was pecked, only the S1 LED was on. For example, when playing song A, pecking sensor S2 caused the food hatch to open while pecking sensor S3 resulted in the pre-set dark time, and vice versa. If the bird did not respond within 25 seconds, a test trial would automatically end without food reward or light-off penalty. Once the accuracy rate of pecking each sensor was greater than 0.60 per day, the duration of the light-off period went from three to one seconds, the food acquisition time from the initial stage of 12 seconds to the later stage of 10 seconds. The duration of this phase varied from bird to bird, with a median value of 456 (IQR 278-655) hours. The proportion of correct responses out of all sounds that each bird

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responded to was calculated on a daily basis as the individual's discrimination rate for the sound stimuli. When a bird learned to associate the two training sounds with the corresponding correct sensors and had reached a discrimination score for the training stimuli greater than 0.75 for three consecutive days (general discrimination score >0.75 , the accuracy rate of each sensor pecking >0.60 for three consecutive days), it was assumed that the bird was able to discriminate the trained song motifs and the training was switched to a transition phase.

Transition phase

During the transition training phase, training stimuli were identical to that in the discrimination training phase, but reinforcement by food reward or light-off period was reduced to occur randomly in 80% (instead of 100%) of trials. In the remaining 20% of trials (with stimuli identical to the training sounds), the subjects were not reinforced with either food or a light-off period. If the bird kept the same level of discrimination as in the training phase for two days, the test phase began. The duration of the transition phase had a median value of 47 (IQR 46-50) hours.

Probe testing phase

In this phase, 16 test stimuli were introduced for 20% of the pecks on S1. Twelve of these were novel stimuli (belonging to either series 1 or series 2). The remaining 4 test stimuli were non-rewarded training sounds used as control and were presented twice as often as the other test stimuli. These test stimuli (non-rewarded training sounds and novel stimuli) were never reinforced and were randomly interspersed between training stimuli. The remaining 80% were training stimuli with reinforcement. Each test sound was presented until it was given 40 trials. This process took two to three weeks, with a median value of 394 (IQR 339-549) hours. After reaching this, the bird was transferred back to its aviary. The order of stimulus presentation was randomized across subjects.

Analysis

To examine whether the two training groups differed in the speed of discrimination learning, we used the total number of trials up to and including the day on which the learning criterion had been reached. A Mann-Whitney-Wilcoxon Test (R Core Team, 2016) was used to detect differences between the two training groups on learning speed (required training trials) since the number of trials didn't follow the normal distribution.

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The reactions to the different test stimuli can be separated into three categories: a ‘correct response’ (i.e., the bird identifies the modified version of training stimulus A as A and the modified version of training stimulus B as B), an ‘incorrect response’ (responding with pecking the sensor for B if the stimulus was a modification of sound A and vice versa), and a ‘nonresponse’ (not pecking a key). For the statistical analyses, we examined the proportion of correct responses as: $\text{Proportion Correct ('PC')} = \text{Count_Correct} / (\text{Count_Correct} + \text{Count_Incorrect} + \text{Count_Nonresponse})$. We found that there was a strong decline in responding to the test stimuli during the test phase: most birds reduced responding to each novel stimulus after 10 presentations (Fig. A1), indicating that the birds apparently learned to recognize the test stimuli as being different from the training ones and providing no reward. For this reason, we restricted our analyses of the responses to the different test stimuli to the first 10 test trials for each stimulus, as during this phase, the responses to the test stimuli were highest and therefore provided the best insights into whether there was variation in the proportion of correct responses between the experimental groups. To examine whether the birds still discriminated the test stimuli above chance, we examined whether the ratio of ‘Count Correct/Count Incorrect’ differed from 1. We did so by applying the $\log(\text{Count_Correct} / \text{Count_Incorrect})$ (indicated as ‘Log(Cor/Inco)’ from now on as the response variable against a $\log(\text{Odds-ratio}) = 0$ in the model analysis. The nested structure of the data was also incorporated into the analysis since, for each experimental group of birds, one half was tested with test stimuli from series 1 and the other with test stimuli from series 2. In addition, one female individual in the Equal-duration training group exhibited responses that significantly deviated from those of the other individuals in the same group. During the probe testing phase, this bird’s proportions of correct responses to two novel versions of stimuli (‘Pitch-shifted +8%’ and ‘Contour-maintained Vcoded’) exceeded 1.5 times the interquartile range (IQR) above the upper quartile (Q3). Consequently, we identified this individual as an outlier and excluded its data from the model analyses (but, for completeness, it is shown in Fig. 4 & Fig. 5).

For the spectrally-changed treatments, the counts of the responses to modified sounds A and B were combined. For the Time-scaled treatments, the ‘PC’ and ‘Log(Cor/Inco)’ were calculated based on the response counts to the stimuli derived from training sound A and those derived from sound B separately. We analysed the data in this way because, for the Unequal-duration group, the ‘Duration stretched 50%’ sound B had a similar duration as Training sound A (sound A was always the longer training sound), and the ‘Duration compressed 50%’ sound

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A had a similar duration as Training sound B. Therefore, we expected that if stimulus duration was a parameter to which the birds were sensitive, that time scaling would differentially affect the responses to changes in the duration of training stimulus A and stimulus B. We thus did a separate analysis for the four Time-scaled treatments ('Duration stretched 20%', 'Duration compressed 20%', 'Duration stretched 50%' & 'Duration compressed 50%') and their corresponding training stimuli for two training groups, comparing the responses to training sounds A and B with those to the Time-scaled versions of sounds A and B.

To investigate the birds' ability to discriminate between various test sounds, Generalized Linear Mixed-effects Models (GLMMs) were utilized. These models incorporated 'Training_Group', 'Test_Treatment' and 'TrainingTrails_scaled' as fixed effects, with 'Bird_ID' as the random effect factor. Additionally, a fixed factor, 'Training_Sound', was included for the Time-scaled test treatments, encompassing four categories: 'Sound A - Equal-duration group', 'Sound B - Equal-duration group', 'Sound A - Unequal-duration group' and 'Sound B - Unequal-duration group'. As 'sex' had a negligible impact at the training group level it was not included in the model analysis. The analysis of these binomial models was carried out in R (R Core Team's methodologies, 2016), utilizing the 'glmer' function from the lme4 package (Bates et al., 2015). Model selection was carried out using the Wald Chi-Squared test. Finally, a post-hoc analysis was conducted on the chosen model, incorporating False Discovery Rate (FDR) correction using the emmeans package (Lenth, 2023).

Ethical Note

The experiment and procedures adhered to the European and Dutch legislation on animal experimentation and were approved by the Dutch Committee for Animal Experimentation (CCD - AVD number 1060020197507) and performed according to the guidelines of the Leiden University Committee for Animal Experimentation

None of the birds had any experience with this experimental setup or the stimuli preceding the experiment. Each experimental bird underwent a physical examination before being transferred to the Skinner boxes. When the birds were in the Boxes, their condition was monitored daily by visual observation throughout all phases of the experiment. The standard checks included: freshwater intake, amount of food obtained in response to pecking sensors, activity, and measuring weight when deemed necessary. The functioning of the operant equipment and stimulus playback were also checked on a daily base. The daily welfare checks

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were done by the experimenter (Ning Z) as well as the qualified animal caretaker (in possession of a so-called ‘art.13f2’ qualification – the qualification which the Dutch law requires), who also advised on the most suitable protocol. Food and water were refreshed three times per week, and the litter floor (containing hard paper and dry sand) of the Skinner Box was cleaned once per week. The food used as reward in the operant chamber consisted of a standard seed mixture for small seed-eating birds (a commercial tropical seed mixture: Deli Nature 56-Foreign finches super, Schoten, Belgium) enriched with mineral and vitamin powder (GistoCal, Raalte, the Netherlands). Cuttlefish bone was also available. This was the same diet as in their home aviary.

If a bird did not operate the sensors for food for more than 18 hours (a very rare event), the hatch would open automatically, allowing a bird to gain sufficient food (approximately equal to the amount of food it should have obtained otherwise), before switching back to the experimental protocol again. The food consumption was checked by recording the amount of food disappearing from the food container. The 18 hours included the 10.5 hours of darkness and meant that the birds would never have been without food for a full day. In addition, obtaining food from the food hatch always gave rise to seeds falling on the floor and this was thus available continuously.

The decision to keep the birds in the Skinner Boxes for the entire duration of the experiment rather than taking the birds in and out of their experimental cage for daily sessions has been discussed with and approved by the Leiden University animal welfare body. The considerations were that daily sessions would require catching and moving the birds – events considered stressful to the birds. Also, in our setup the birds could get access to food whenever they wanted whereas otherwise some food restriction period would be necessary to keep the birds motivated. The training stimuli were normal zebra finch songs, which are known to be attractive to both male and female zebra finches. After finishing all phases of the experiment, the birds were returned to their home aviaries. Previous similar experiments showed that birds reintroduced to the aviaries after having been in the Skinner Boxes for several weeks experienced no particular difficulties.

RESULTS

Speed of Discrimination Learning

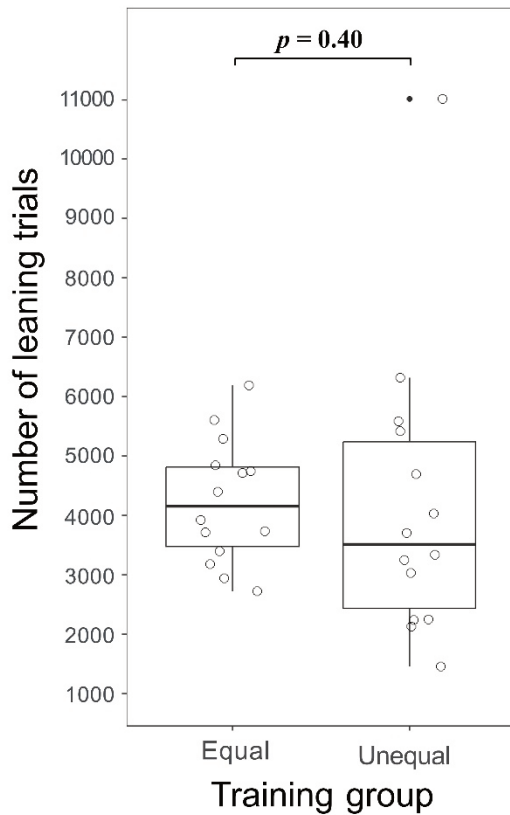


Figure 4. Number of learning trials needed to reach the learning criterion. Individual zebra finch results are shown with open dots. There is no significant difference between the Equal-duration group and the Unequal-duration group. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range.

The discrimination training lasted until a bird reached the learning criterion of over 75% correct responses to both sound A and sound B for consistent three days. All 28 birds finished the training and learned the discrimination on an average of 4209 (SD = 1840, N = 28) trials to reach the criterion. No significant difference ($p = 0.40$, $z = 0.87$; Fig. 4) was found between the Equal-duration group (M = 4243, SD = 1041) and the Unequal-duration group (M = 4175, SD = 2439). Removal of one outlier (a female individual from the Unequal-duration training group requiring 11011 learning trials) did not change this outcome. It suggests that birds from two training groups learn approximately equally fast.

The responses to test stimuli

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We examined the impact of the stimulus modifications in several ways. First, we examined whether spectral changes (frequency shifts as well as vocoding) had an impact on the proportion of correct identifications of the stimuli. Doing so, we address whether this impact was in the predicted direction of being larger in the Equal-duration than in the Unequal-duration group, based on the assumption that spectral changes might serve as primary cues in the Equal-duration group, given the (almost) identical song durations. Next, we examined whether there is a difference in impact among the various spectral modifications. However, while the proportion of correct responses may be affected by a modification, this need not imply that the birds can no longer discriminate between similar modifications of training songs A and B; they may still show more correct than incorrect responses. To address this, we examined whether the ratio of correct *versus* incorrect responses to a modified stimulus was still above chance. Finally, we use the same structure to examine the impact of the tempo changes on the birds' proportions of correct responses and discrimination rate.

The effect of spectral changes

Responses to spectrally-changed stimuli differ between groups and between test stimuli

For the birds' responses to stimuli that are spectrally manipulated, the ANOVA Type III for models of both test series showed that the proportion of correct responses (PC) differed significantly between the Equal-duration and Unequal-duration training groups as well as among the different Test stimuli. Thus, the two factors 'Training_Group' and 'Test_Treatment', as well as their strong interaction effects for the response variable 'PC' were selected as fixed factors for models of both series (see ① and ② in Table 2), in addition, the factor 'TrainingTrails_scaled' and its interaction with 'Test_Treatment' were left in the model for the response variable 'PC' in series 1 since a significant effect was found in this interaction (see ① in Table 2).

Table 2 ANOVA (Type III Wald chi square tests) table for selected GLMs

Variable	Chisq.	Df	Pr(>Chisq.)
① MODEL PC for spectrally-changed (Sound A & Sound B combined in Series 1)			
Training_Group	4.0689	1	0.0437 *
Test_Treatment	95.1957	4	< 2.2e-16 ***
Training_Group:Test_Treatment	11.4205	4	0.0222 *

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TrainingTrails_scaled	0.0161	1	0.8990
Test_Treatment:TrainingTrails_scaled	14.1089	4	0.0070 **
② MODEL PC for spectrally-changed (Sound A & Sound B combined in Series 2)			
Training_Group	9.3598	1	0.0022 **
Test_Treatment	101.5756	4	< 2.2e-16 ***
Training_Group:Test_Treatment	9.7748	4	0.0444 *
③ MODEL PC for time-scaled (Sound A & Sound B separated in Series 1)			
Training_Group	0.0053	1	0.9421
Test_Treatment	48.2401	2	3.348e-11 ***
Training_Sound	0.0633	2	0.9689
Training_Group:Test_Treatment	9.0091	2	0.0111 *
Test_Treatment:Training_Sound	10.2016	4	0.0372 *
④ MODEL PC for time-scaled (Sound A & Sound B separated in Series 2)			
Training_Group	1.2867	1	0.25666
Test_Treatment	13.9044	2	0.00096 ***
Training_Sound	0.5413	2	0.76290
Training_Group:Test_Treatment	0.3930	2	0.82159
Test_Treatment:Training_Sound	2.4069	4	0.6613

Note: All variables shown here were used as fixed factors for corresponding models, no matter their p values were significant or not because these were the variables of our interest. ‘Bird_ID’ was used as the only random factor in all models. • $0.05 < P \leq 0.10$, * $0.01 < P \leq 0.05$, ** $0.001 < P \leq 0.01$, *** $P \leq 0.001$.

Spectral changes affect the Equal- duration group most strongly

Figure 5 shows that the Equal-duration group had a lower PC to all spectrally-changed test stimuli compared to the Unequal-duration group. The pairwise comparisons between two training groups by the Post hoc Tukey’s HSD tests (Table A1) showed that this difference was significant for the stimuli ‘Pitch-shifted -20%’ ($P < 0.01$) and ‘Contour-averaged Vcoded’ ($P < 0.05$) in series 1 and for the stimuli ‘Pitch-shifted -8%’ ($P < 0.05$), ‘Frequency-shifted 500Hz’ and ‘Contour-maintained Vcoded’ (both $P < 0.01$) in series 2. In addition, Pitch-shifted upward versions seem to have less impact on the between-groups difference than Pitch-shifted downward versions.

The observed differences between the groups are in line with our expectation that birds trained with Equal-duration stimuli are more sensitive to spectral changes than the birds trained with Unequal-duration stimuli. They also show that this effect is present in both test series.

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Differences in responses among the test stimuli

In both series 1 and series 2, the birds responded with a higher PC to the training stimuli compared to all four spectrally-changed stimuli in both training groups. For each training group, we examined whether there were differences in the PC of birds' responses among four spectrally-changed stimuli for each test series (Table A1). In series 1 the birds of the Equal-duration group responded with a significantly higher PC to the 'Pitch-shifted +20%' stimulus than to 'Contour-averaged Vocoded' ($P < 0.05$), and a clear trend to a difference between 'Frequency-shifted 1500Hz' and 'Contour-averaged Vocoded' ($P = 0.06$). The birds of the Unequal-duration group responded with a significantly higher PC to the 'Pitch-shifted -20%' stimulus than to 'Contour-averaged Vocoded' and 'Frequency-shifted 1500Hz' (both $P < 0.05$) (Fig. 5a). In series 2, the birds of the Equal-duration group responded with a significantly higher PC to the 'Pitch-shifted +8%' stimulus than to the other three spectrally-changed stimuli ('Pitch-shifted -8%' ($P < 0.05$), 'Frequency-shifted 500Hz' ($P < 0.01$) & 'Contour-maintained Vocoded' ($P < 0.001$)). For the Unequal-duration group there is no significant difference in PC among the four spectrally-changed stimuli (Fig. 5b). On the whole, these results show a weak tendency that pitch-shifted versions have less impact on the PCs than vocoded versions. This implies that, if anything, the zebra finches are attending more strongly to precise spectral details of the song elements rather than to the spectral envelope. If they would attend more to the latter, vocoding would have a lesser impact than the other manipulations.

Are spectrally-changed stimuli still recognized?

If the birds are still capable of linking the modified stimuli to the respective training stimuli, the number of correct responses to the test stimuli should be higher than the number of incorrect responses. The birds of the Unequal-duration group responded above chance to all spectrally-changed stimuli in both two test-series (Fig. 6a, b), while birds of the Equal-duration group responded above chance only to two of the spectrally-changed stimuli ('Pitch-shifted +8%' & 'Frequency-shifted 500Hz') in series 2, and to none of the spectrally-changed stimuli in series 1 (Table A2). This confirms the finding above that the birds from the Equal-duration group are more strongly attending to spectral features than the birds from the Unequal-duration group. In addition, the Unequal group showed a lower degree of recognition when the modifications were stronger (series 1) than when they were less strong (series 2).

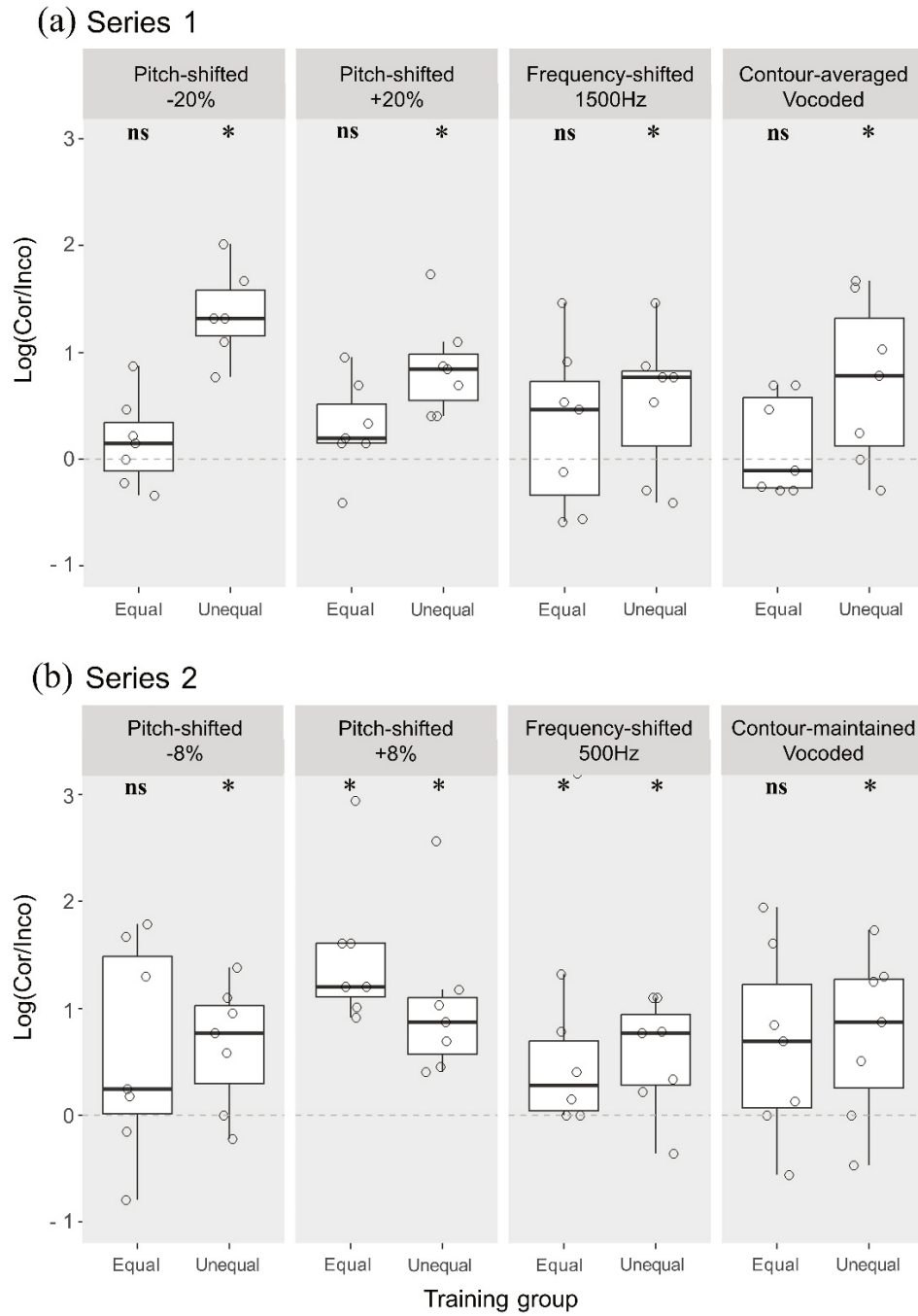
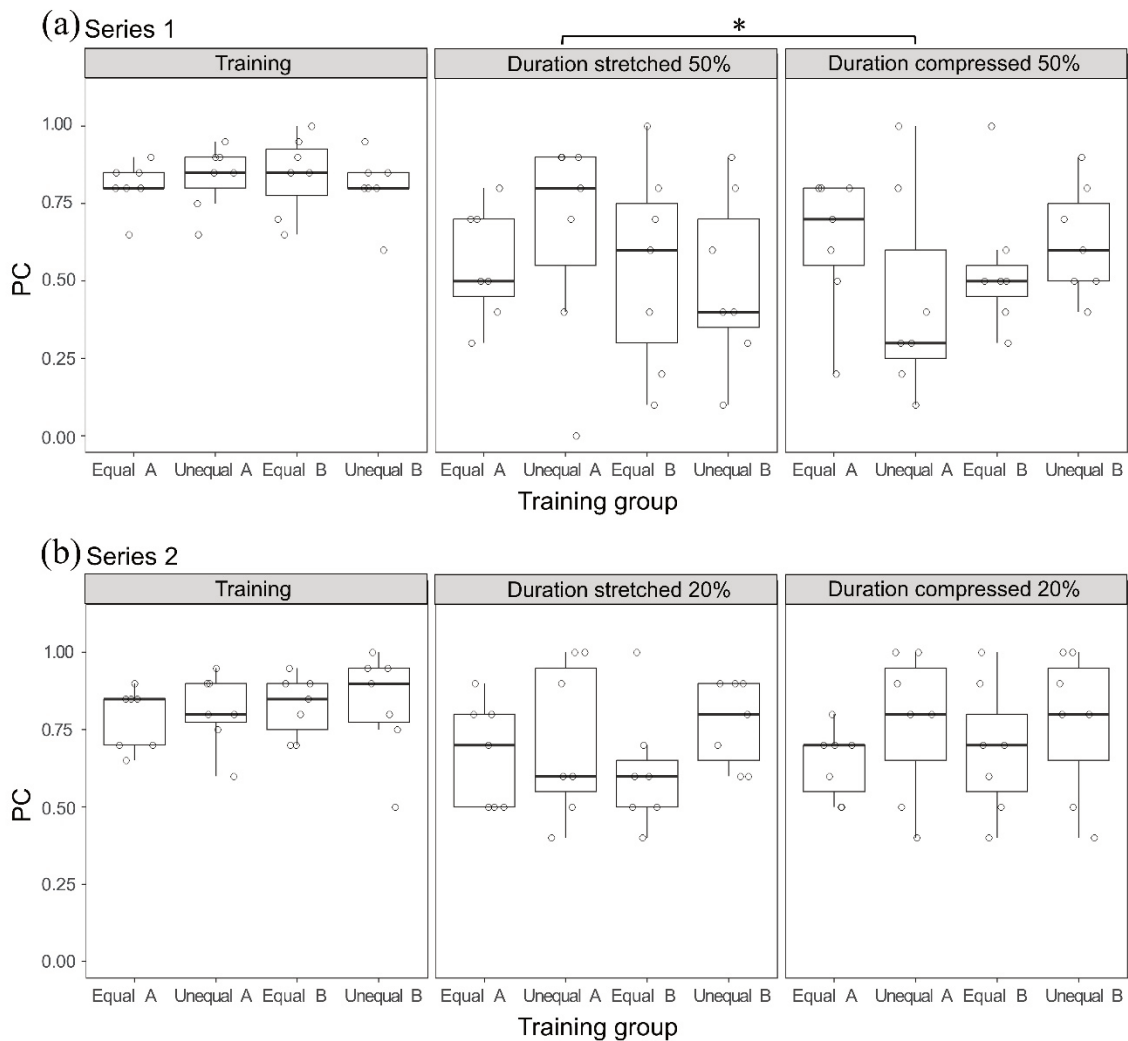


Figure 6. Visualisation of $\text{Log}(\text{Cor}/\text{Inco})$ for birds responding to spectrally-changed stimuli of series 1 (a) and series 2 (b). A * indicates that the $\text{Log}(\text{Cor}/\text{Inco})$ of a Test treatment is significantly different from 0; ns indicates that the $\text{Log}(\text{Cor}/\text{Inco})$ of a Test treatment is overlapping with 0. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range. Horizontal dashed lines show the discrimination boundaries in which the proportion of correct responses is equal to the proportion of incorrect responses. The calculation of $\text{Log}(\text{Cor}/\text{Inco})$ s was based on the counts of ‘correct response’ and ‘incorrect response’ from the same data set that was also used for Fig.5. Note that one bird’s data point cannot be fully displayed on the panel (b) because it made no incorrect responses to the ‘Frequency-shifted 500Hz’ stimuli version, resulting in an infinitely large value after log-scaling.

The effect of duration changes

Tempo changes affect the Equal- and Unequal-duration group differently

In the ANOVA Type III model for responding to Time-scaled stimuli we also included the factors ‘Training_Sound’ (A or B) as fixed factor in addition to the factors ‘Training_Group’ and ‘Test_Treatment’, as well as the interactions of ‘Training_Group’ and ‘Training_Sound’ with ‘Test_Treatment’. There were no significant differences in PC between the Equal-duration and the Unequal-duration training groups if the results for the Time-scaled versions of training stimuli A and B are combined. However, the results showed a significant interaction effect between Training Group and Training Sound, as well as for Test Treatment and Training Sound for series 1 (see ③ in Table 2). Figure 7 shows that this is due to the different responses of both groups to the various Time-scaled versions of training sounds A and B.



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Figure 7. Proportion correct (PC) of responses to Time-scaled stimuli of series 1 (a) and series 2 (b).). For significant differences between Training and Duration-changed stimuli: see text. The only difference among the Duration-changed stimuli was present in series 1: * refers to a significant difference of $0.01 < P \leq 0.05$. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range.

Differences in responses among the test stimuli

The pairwise comparisons of the PC for the Time-scaled versions of sound A and sound B for the Equal- and Unequal-duration groups are shown in Table A3. In both series 1 and series 2, there is no significant difference in the PC between all three sound versions (Training and two Time-scaled versions) between UnequalA and UnequalB, and between EqualA and EqualB. However, in series 1, the birds responded with a higher PC to the Training stimuli compared to the Time-scaled versions of soundA and soundB (in both Equal-duration and Unequal-duration groups). This difference is significant in all comparisons apart from the difference between the training A and the ‘Duration compressed 50%’ A in the Equal-duration group, which showed a clear trend in the same direction ($P = 0.06$). The birds of the Unequal-duration group responded with a significantly lower PC to the ‘Duration compressed 50%’ A than to the ‘Duration stretched 50%’ A ($P < 0.05$) (Fig. 7a).

In series 2, the birds of the Equal-duration group responded with a significantly higher PC to Training B than to ‘Duration stretched 20%’ B ($P < 0.01$), and a clear trend of difference between Training B and ‘Duration compressed 20%’ B ($P = 0.06$). For the Unequal-duration group the PC did not differ between the Training sounds and the 20% Duration changed stimuli. To conclude, the ‘ $\pm 50\%$ ’ Time-scaled manipulation was noticed by birds from both Equal-duration and Unequal-duration groups, and such an impact was weaker in the ‘ $\pm 20\%$ ’ Time-scaled manipulation. In addition, the Unequal-duration group responded differently to whether the 50% duration change concerned the long or the short song. This difference is meaningful and was expected if the birds in the Unequal-duration group attend to the song duration for song recognition. Training sound A was always 50% longer than training sound B in the Unequal-duration training group. Therefore the ‘Duration compressed 50%’ of the sound A stimulus made this stimulus the same length as training stimulus B, while the ‘Duration stretched 50%’ of the sound B stimulus made this stimulus the same length as training stimulus A. This suggests that the similarities in duration between training songs and test songs resulted in reduced song recognition even when there were still differences in spectral features between the pair of sounds.

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Are time-scaled stimuli still recognized?

At the group level (if the birds' responses to sound A or sound B are not differentiated in the analysis), the responses of birds of both groups to all stimuli (Training stimuli and Time-scaled versions) are all different from 0, indicating they are recognized (Table A4). Similar to the analysis for the spectrally-changed stimuli, we also examined for which of the Time-scaled stimuli the number of correct responses was higher than that of the incorrect responses, but now differentiating between the responses to the test stimuli derived from training sound A and those derived from B. The birds of the Equal-duration group responded correctly above chance on the 'Duration stretched 50%' and 'Duration compressed 50%' sound A, but not to the 'Duration stretched 50%' and 'Duration compressed 50%' sound B (Fig. 8a). However, as for this group training songs A and B were of Equal-duration duration and arbitrarily assigned to be either A or B, the difference between the responses can be ascribed to chance, also because there is no significant difference between the scores to the variants derived from training stimulus A and B (Table A5). The birds of the Unequal-duration group respond significantly above the chance to the 'Duration stretched 50%' sound A and 'Duration compressed 50%' sound B but respond to the 'Duration stretched 50%' sound B and 'Duration compressed 50%' sound A by chance (Fig. 8a). In line with the finding of a difference in impact on the proportion of correct responses, the difference between recognizing the stretched and the compressed versions of sounds A and B by birds from the Unequal-duration group confirms that these birds used song duration to distinguish training songs A and B. In series 2, the responses of birds of both groups to all Time-scaled sounds (no matter whether it was sound A or sound B) are statistically different from 0 in favour of correct response (Fig. 8b).

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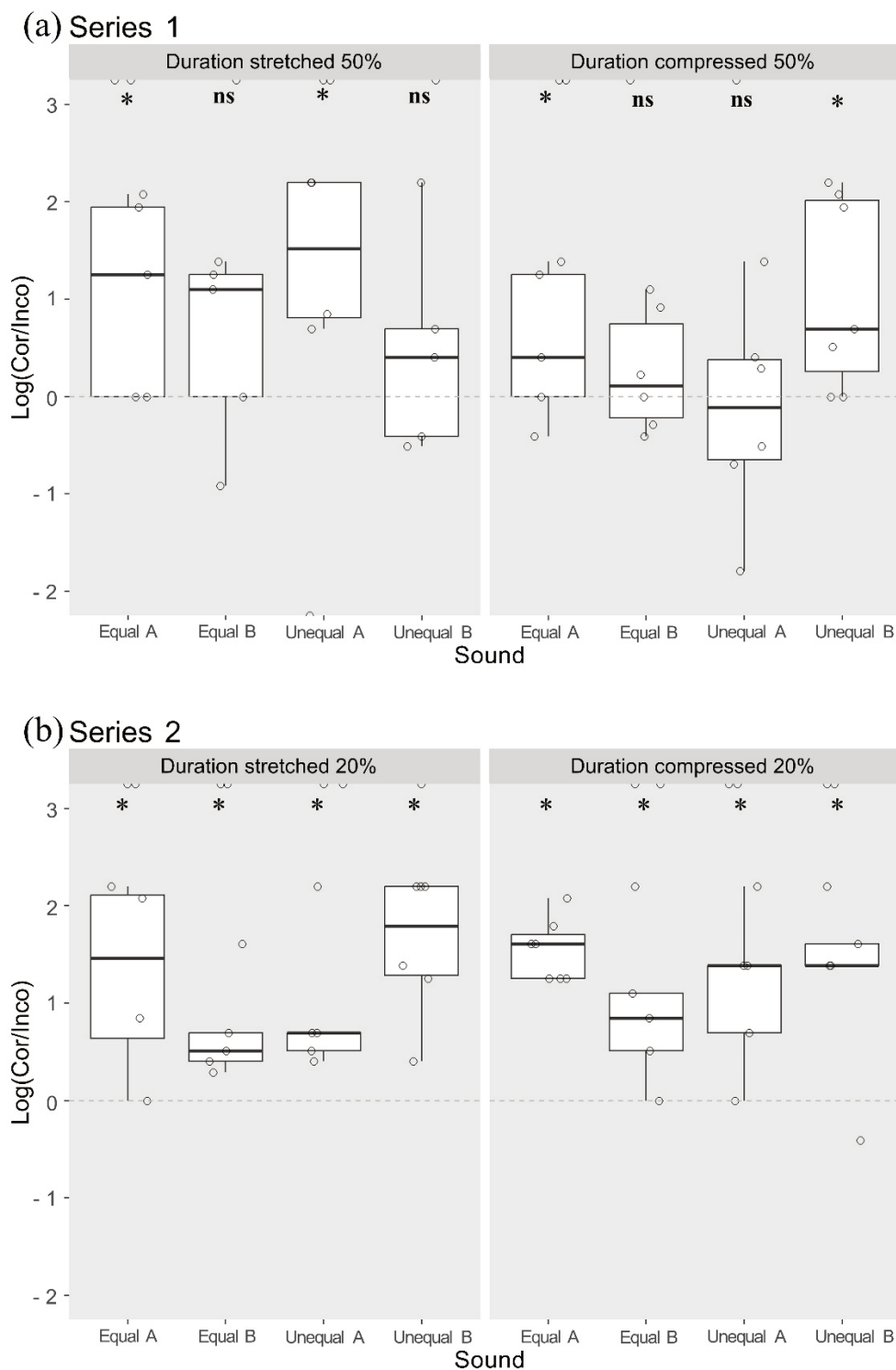


Figure 8. Visualisation of Log (Correct/Incorrect) for birds responding to time-scaled stimuli of series 1 (a) and series 2 (b). A * indicates that the Log(Cor/Inco) of a Test treatment is significantly different from 0; ns indicates that the Log(Cor/Inco) of a Test treatment is overlapping with 0. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range. Horizontal dashed lines show the discrimination boundaries in which the proportion of correct responses is equal to the proportion of incorrect responses. The calculation of Log(Cor/Inco)s was based on the counts of ‘correct response’ and ‘incorrect response’ from the same data set that was also used for Fig.7.

DISCUSSION

Our study demonstrates that zebra finches can use both spectral features and song duration when discriminating between two songs. However, the importance of these acoustic parameters depended on whether the songs differed in duration or not, with spectral features having a less prominent role when duration is available as an additional feature to distinguish two songs. Our results thus show that the acoustic parameters that zebra finches attend to are, at least partially, context-driven, i.e., dependent on the degree to which these parameters differ between songs and as such support the hypothesis that zebra finches are cognitively flexible in their attention to different acoustic parameters, related to the salience of the differences between songs.

Song duration does not affect learning speed

If the zebra finches can use song duration as an additional cue for discrimination learning, then we may expect that this results in faster song discrimination learning with songs of different compared to similar duration. However, in our current experiment, the learning speed of the birds trained on songs of unequal duration does not differ from the birds trained on songs of equal duration. Combined with our test results indicating that both experimental groups attend to spectral cues as well as song duration, albeit with a difference in weight, this suggests that both song features are considered right from the start of the learning process.

Spectral cues or song duration?

Various studies (Dooling et al., 2002; Lohr et al., 2006; Vernaleo et al., 2010; Vernaleo & Dooling, 2011; Lawson et al., 2018; Prior et al., 2018a; Prior et al., 2018b; Geberzahn & Derégnaucourt, 2020; Fishbein et al. 2021; Mol et al., 2021) concluded that when zebra finches discriminate two songs they primarily attend to spectral details and temporal fine structure within individual syllables, and are far less sensitive to syllable sequence and temporal features of the whole song, such as song duration. In particular, a study by Nagel et al. (2010) showed that an 8% pitch-shift already resulted in reduced discrimination between two songs, and that the songs were no longer discriminated after a 32% pitch-shift. In contrast, stretching or compressing the songs by 32% in duration hardly affected discrimination, which was

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maintained even after a 64% change. In that study zebra finches had to discriminate two songs of equal duration, hence the birds could not use song duration to recognize the songs. However, several recent studies indicated that the parameters, that zebra finches can or do use in discriminating and recognizing sounds, may depend, at least to some extent, on the difference between the sound stimuli (Burgering et al., 2018; Burgering et al., 2019; Ning et al., 2023). Therefore, the main question underlying the current experiment was whether the importance of spectral parameters ('relative pitch' and 'spectral envelope'), and the temporal parameter 'duration', depended on whether the songs that had to be discriminated, differed in overall duration. In both test series, the birds from both the Equal- and the Unequal-duration groups responded with a lower proportion of correct responses to all four spectrally-changed stimuli than to the training stimuli, indicating that all birds were able to detect all the different types of spectral changes. However, the impact of the spectral changes was stronger in the Equal-duration than in the Unequal-duration group for both test series. The impact of the spectral changes was also stronger in series 1, in which the test sounds featured more substantial spectral modifications compared to the training sounds than in series 2. For the Equal-duration group, this even resulted in a loss of recognition of spectrally modified versions of training sounds for all spectral modifications in series 1 and half of them in series 2, while the Unequal-duration group maintained the recognition of all spectrally modified stimuli in both series. In response to changes in song duration, both groups also showed a lower proportion of correct responses and poorer discrimination when song durations were stretched or compressed by 50% (series 1), thus indicating that both groups attended to song duration. However, a 20% change in duration showed only a limited effect. These results are within the same ranges as observed by Nagel et al. (2010). Nevertheless, the importance of song duration for song discrimination was very noticeable in the Unequal-duration group. These birds no longer discriminated between the songs when the 50% compressed and stretched versions made the test song of the same length as the opposite training song, i.e., when the duration of the manipulated song A is similar to the duration of training song B and vice versa. For the Unequal-duration group, the 20% Time-scaled manipulation affected the discrimination substantially less than the 50% Time-scaled manipulation. In this case, the temporal manipulation did not eliminate the differences in song duration between manipulated and training songs. To conclude, while our study confirms the important contribution of spectral features for song discrimination as obtained in earlier studies, it also shows that zebra finches use song duration as a prominent feature when songs are substantially different in duration, at the expense of attending to

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spectral features. Future studies may address whether the impact song duration is related to the magnitude of the difference in duration between songs.

The finding that zebra finches are attending to the absolute duration of a stimulus also has relevance for studies examining rhythm perception in zebra finches. The crucial test for being able to perceive a rhythmic pattern is whether humans or animals can recognize a melody or tone sequence when this sequence is being speeded up or slowed down (e.g., Bouwer et al., 2021). Several studies demonstrated that zebra finches could discriminate between a regular and an irregular pattern of song syllables or artificial tones (Lampen et al., 2014; van der Aa et al., 2015; ten Cate et al., 2016; Lampen et al., 2017; Rouse et al., 2021, 2023). However, this discrimination is reduced with a tempo change of the stimuli (van der Aa et al., 2015; ten Cate et al., 2016). This indicates that zebra finches attend more to the absolute duration of components of a stimulus, such as the duration of specific elements or intervals, rather than to the overall pattern of regularity (ten Cate et al., 2016), although it might be that with extensive training zebra finches might become more sensitive to the overall pattern (Rouse et al., 2021; 2023). The current finding that zebra finches show reduced discrimination when songs are compressed or stretched, and attend to absolute song duration thus is in line with the results of the studies on zebra finch rhythm perception.

Impact of various spectral changes

The second question we aimed to address in the current study concerns the relevance of spectral envelope and pitch in song discrimination. All our spectral manipulations maintained the absolute durations of syllables and songs but affected the spectral structure in different ways. The Frequency-shifted test stimuli moved the whole spectrum upward in a linear way. This maintained the frequency bandwidth but changed the harmonic relationships (with harmonic overtones being converted into inharmonic partials) among the frequencies within and between syllables. In the pitch-shifted stimuli the relative relationships among the frequencies within and between syllables are maintained, but the absolute pitch values have changed from those of the training stimuli. For the vocoded stimuli, the frequency ranges (spectral envelope) are identical to the training stimuli, but pitch information is removed and replaced by noise. Although the Unequal-duration group used the duration as a prominent cue and was less affected by the spectral changes, both groups showed decreased discrimination of all spectrally changed stimuli compared to discrimination of the training stimuli. Overall,

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the vocoded versions seem to reduce the discrimination more than the other stimuli, with at best a weak tendency that discrimination is maintained best for the pitch-shifted stimuli. If we compare our data on the impact of pitch shifts on song discrimination with those obtained by Nagel et al. (2010), we found that birds in the Equal-duration group, which is most comparable to the experiment by Nagel et al. (2010), show a comparable outcome. In that study, an 8% pitch shift reduced but still maintained discrimination, but a 32% shift resulted in a lack of discrimination. These effects are in the same range as the reduced discrimination we obtained with an 8% pitch shift and lack of discrimination with a 20% change. The results of both our study and that by Nagel et al. (2010) also indicate that zebra finches are more sensitive to pitch changes of songs than starlings are, which can still show discrimination of songs with pitch shifts up to $\pm 40\%$ (Bregman et al., 2012). Interestingly, starlings trained on piano melodies responded more strongly to pitch changes than those trained on songs, indicating that the nature of the stimuli may be a relevant factor in this songbird's sound discrimination (Bregman et al., 2012).

Finally, we showed that both types of vocoded stimuli strongly reduced discrimination of the songs to a similar extent. It thus did not matter whether the spectral contour was maintained over the elements (Contour-maintained Vocoded) or not (Contour-averaged Vocoded). The impact of noise-vocoding on song recognition is surprising in light of earlier studies. For starlings, Bregman et al. (2016) showed that vocoded versions, but not pitch-shifted versions, of sequences of tones that varied in pitch and timbre, maintained the discrimination between these sequences. This indicated that the sequences were discriminated by their spectral envelope rather than pitch. Patel (2017) argued that this might be a common characteristic across birds, also for the discrimination of natural vocalizations. However, so far, no study examined how starlings respond to vocoded versions of conspecific songs and it hence remains to be explored whether such a stimulus would result in similar outcomes when compared to testing with artificial sounds. Nevertheless, the importance of the spectral envelope for auditory discrimination in birds seemed to be supported by a study in zebra finches, in which Burgering et al. (2019) trained zebra finches to distinguish two sets of artificial harmonic tone stimuli, which could only be differentiated by attending to the spectral envelope. When these stimuli were noise-vocoded, maintaining the (absolute) spectral envelope but removing (absolute) pitch information, the discrimination was maintained, indicating that zebra finches indeed attended to the spectral envelope of the stimuli. Also, an extensive analysis of zebra finch vocalizations indicated that the shape of the frequency spectrum (spectral envelope) of

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the different vocalizations was an important potential information-bearing feature (Elie & Theunissen, 2016) for distinguishing various vocalizations. Hence, one would expect spectral envelope to be important for discriminating songs. Why this does not show up in the current study is not clear. One factor might be that the spectral envelope might be relevant to zebra finches when discriminating among calls or other shorter sounds, such as the single element stimuli used by Burgering et al. (2019). In contrast, discrimination of songs might rely more on attending to other spectral features, including pitch and harmonic structure of the songs. Attendance to such features has been demonstrated in a range of studies (e.g., Okanoya & Dooling, 1990; Uno et al., 1997; Dooling & Lohr, 2006; Lohr et al., 2006; Vignal & Mathevon, 2011; Prior et al., 2018a; Prior et al., 2018b).

Conclusion and outlook

To conclude, our study shows that the acoustic parameters that zebra finches use to distinguish between different songs depend on the dimensions in which these songs differ. As we demonstrate here, this could be spectral features, but also song duration. Similarly, in another study we showed that although zebra finches usually do not give much attention to the sequential order of the syllables when discriminating songs, they can use this sequence quite readily if needed (Ning et al., 2023), while Burgering et al. (2018, 2019) demonstrated that also the attendance to either the fundamental frequency (absolute pitch) or the energy distribution of a harmonic spectrum (spectral envelope) varied depending on the task. These results thus contribute to expanding evidence that zebra finches are cognitively flexible: when faced with the task of discriminating between different acoustic stimuli, they appear to focus on the most salient features distinguishing these stimuli. That the importance of different parameters for sound discrimination may depend on the nature of the stimuli and on the task the birds facing is also recognized by others (e.g., Patel, 2017). However, this does not imply that there is no bias in this ability, but it indicates that there may be a difference between which features an animal *does* use to discriminate stimuli in a particular context and which it *can* use. Our study also shows that both the features in which stimuli differ, as well as the magnitude of those differences, affect their importance in discrimination. Future studies might explore other potential cues for song discrimination. Such investigations will contribute to a more nuanced understanding of how birds perceive and utilize various song features as a discriminative cue. At the same time, comparing the results of our study with those obtained

in starlings (Bregman et al., 2016) suggests important differences between avian species, differences that call for further exploration.

Acknowledgments

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Appendix Figures & Tables

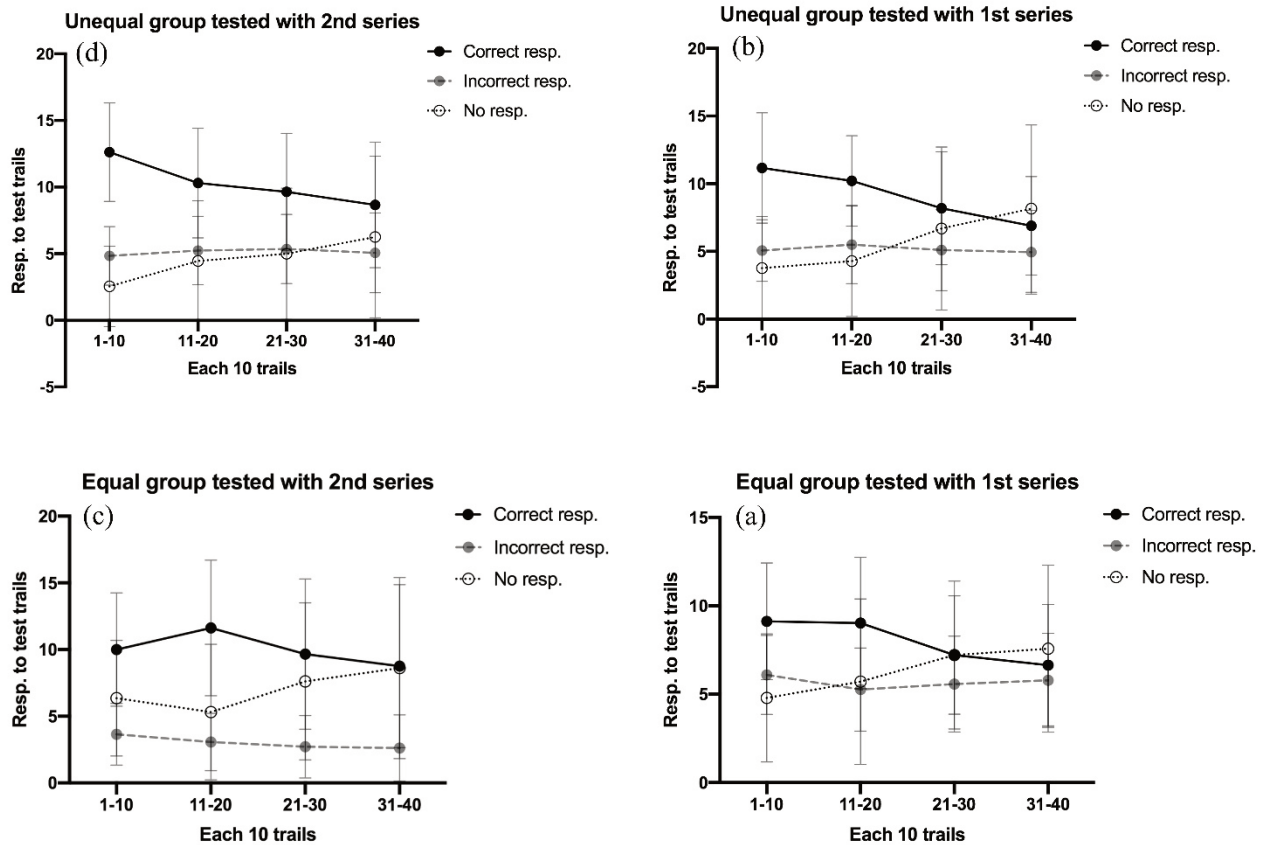


Figure A1. Counts of birds' responses to the test stimuli during the test phase. (a) Trials of Equal-duration group and (b) Unequal-duration group responding to 1st series of test stimuli; (c) Trials of Equal-duration group and (d) Unequal-duration group responding to 2nd series of test stimuli. The total 40 test trials were divided into four 10-trials blocks. Lines across four 10-trials blocks refer to three categories of reaction to a sound: the 'correct response' (Black line), the 'incorrect response' (Black dash line), and a 'nonresponse' (Grey dash line).

Table A1 Post hoc test results of Binomial GLM for the interaction of spectrally-changed Test_Treatment & Training_Group

Stimuli	Training_Group	Series	estimate	SE	Z.ratio	P-value
a. PC (sound A+B in two training groups)						
Pair wise comparisons between training groups - Series 1						
Training	Equal - Unequal	1	0.0676	0.407	0.166	0.8680
Pitch-shifted -20%	Equal - Unequal	1	-1.1345	0.359	-3.165	0.0080
Pitch-shifted +20%	Equal - Unequal	1	-0.6006	0.351	-1.711	0.1450
Frequency-shifted 1500Hz	Equal - Unequal	1	-0.2342	0.349	-0.671	0.6281
Contour-averaged Vocoded	Equal - Unequal	1	-0.8465	0.355	-2.387	0.0425
Pair wise comparisons between test stimuli - Series 1						
Training - Pitch-shifted -20%	Equal_Duration	1	2.1898	0.294	7.456	<.0001
Training - Pitch-shifted +20%	Equal_Duration	1	1.8516	0.290	6.384	<.0001
Training - Frequency-shifted 1500Hz	Equal_Duration	1	1.9422	0.290	6.688	<.0001
Training – Contour-averaged Vocoded	Equal_Duration	1	2.4698	0.298	8.301	<.0001
Pitch-shifted -20% - Pitch-shifted +20%	Equal_Duration	1	-0.3382	0.250	-1.352	0.2519
Pitch-shifted -20% - Frequency-shifted 1500Hz	Equal_Duration	1	-0.2476	0.250	-0.989	0.3585
Pitch-shifted -20% – Contour-averaged Vocoded	Equal_Duration	1	0.2800	0.258	1.084	0.3478
Pitch-shifted +20% - Frequency-shifted 1500Hz	Equal_Duration	1	0.0906	0.246	0.368	0.7128
Pitch-shifted +20% – Contour-averaged Vocoded	Equal_Duration	1	0.6182	0.254	2.431	0.0301
Frequency-shifted 1500Hz – Contour-averaged Vocoded	Equal_Duration	1	0.5276	0.254	2.074	0.0635
Training - Pitch-shifted -20%	Unequal_Duration	1	0.9877	0.296	3.341	0.0021
Training - Pitch-shifted +20%	Unequal_Duration	1	1.1834	0.292	4.059	0.0002
Training - Frequency-shifted 1500Hz	Unequal_Duration	1	1.6404	0.290	5.652	<.0001
Training - Contour-averaged Vocoded	Unequal_Duration	1	1.5557	0.289	5.374	<.0001
Pitch-shifted -20% - Pitch-shifted +20%	Unequal_Duration	1	0.1957	0.259	0.755	0.5001
Pitch-shifted -20% - Frequency-shifted 1500Hz	Unequal_Duration	1	0.6527	0.257	2.537	0.0224
Pitch-shifted -20% – Contour-averaged Vocoded	Unequal_Duration	1	0.5680	0.256	2.215	0.0446
Pitch-shifted +20% - Frequency-shifted 1500Hz	Unequal_Duration	1	0.4570	0.253	1.810	0.1005

Pitch-shifted +20% – Contour-averaged Vocoded	Unequal_Duration	1	0.3724	0.252	1.480	0.1738
Frequency-shifted 1500Hz – Contour-averaged Vocoded	Unequal_Duration	1	-0.0846	0.249	-0.339	0.7344
Pair wise comparisons between training groups - Series 2						
Training	Equal - Unequal	2	-0.252	0.387	-0.652	0.5146
Pitch-shifted -8%	Equal - Unequal	2	-0.889	0.339	-2.623	0.0145
Pitch-shifted +8%	Equal - Unequal	2	-0.358	0.335	-1.069	0.3565
Frequency-shifted 500Hz	Equal - Unequal	2	-1.095	0.344	-3.187	0.0035
Contour-maintained Vocoded	Equal - Unequal	2	-1.230	0.348	-3.539	0.0020
Pair wise comparisons between test stimuli - Series 2						
Training - Pitch-shifted -8%	Equal_Duration	2	1.9396	0.296	6.545	<.0001
Training - Pitch-shifted +8%	Equal_Duration	2	1.3160	0.291	4.518	<.0001
Training - Frequency-shifted 500Hz	Equal_Duration	2	2.2073	0.302	7.306	<.0001
Training – Contour-maintained Vocoded	Equal_Duration	2	2.3733	0.307	7.731	<.0001
Pitch-shifted -8% - Pitch-shifted +8%	Equal_Duration	2	-0.6236	0.265	-2.350	0.0268
Pitch-shifted -8% - Frequency-shifted 500Hz	Equal_Duration	2	0.2677	0.277	0.966	0.3710
Pitch-shifted -8% – Contour-maintained Vocoded	Equal_Duration	2	0.4337	0.282	1.537	0.1555
Pitch-shifted +8% - Frequency-shifted 500Hz	Equal_Duration	2	0.8912	0.272	3.279	0.0017
Pitch-shifted +8% – Contour-maintained Vocoded	Equal_Duration	2	1.0572	0.277	3.815	0.0003
Frequency-shifted 500Hz – Contour-maintained Vocoded	Equal_Duration	2	0.1660	0.288	0.576	0.5647
Training - Pitch-shifted -8%	Unequal_Duration	2	1.3031	0.286	4.548	<.0001
Training - Pitch-shifted +8%	Unequal_Duration	2	1.2099	0.287	4.214	0.0001
Training - Frequency-shifted 500Hz	Unequal_Duration	2	1.3646	0.286	4.768	<.0001
Training – Contour-maintained Vocoded	Unequal_Duration	2	1.3952	0.286	4.877	<.0001
Pitch-shifted -8% - Pitch-shifted +8%	Unequal_Duration	2	-0.0932	0.249	-0.374	0.8876
Pitch-shifted -8% - Frequency-shifted 500Hz	Unequal_Duration	2	0.0615	0.248	0.248	0.8934
Pitch-shifted -8% – Contour-maintained Vocoded	Unequal_Duration	2	0.0921	0.248	0.372	0.8876
Pitch-shifted +8% – Frequency-shifted 500Hz	Unequal_Duration	2	0.1547	0.249	0.622	0.8876
Pitch-shifted +8% – Contour-maintained Vocoded	Unequal_Duration	2	0.1853	0.249	0.745	0.8876
Frequency-shifted 500Hz – Contour-maintained Vocoded	Unequal_Duration	2	0.0306	0.247	0.124	0.9016

Note: Response variables in GLMMs: the PC of responses if birds respond to one of two sounds. Each of the corrected pairwise multiple comparison tests is separated into clusters within the table. Bold indicates significant differences.

Table A2 Estimates and 95% confidence intervals for the correct identification of spectrally-changed stimuli

Training_Group	Stimuli	Series	Emmean	SE	CL (95%)	
					Lower	Upper
Log(Corr/Inco) ~ Training_Group + Stimuli + Training_Group:Stimuli + (1 BirdID), (Sound A & Sound B combined)						
Pair wise comparisons - Series 1						
Equal_Duration	Pitch-shifted -20%	1	0.201	0.202	-0.1958	0.5976
Equal_Duration	Pitch-shifted +20%	1	0.283	0.191	-0.0908	0.6564
Equal_Duration	Frequency-shifted 1500Hz	1	0.310	0.200	-0.0813	0.7021
Equal_Duration	Contour-averaged Vocoded	1	0.002	0.215	-0.4196	0.4237
Unequal_Duration	Pitch-shifted -20%	1	1.075	0.213	0.6573	1.4921
Unequal_Duration	Pitch-shifted +20%	1	0.833	0.202	0.4373	1.2284
Unequal_Duration	Frequency-shifted 1500Hz	1	0.635	0.206	0.2314	1.0380
Unequal_Duration	Contour-averaged Vocoded	1	0.693	0.205	0.2900	1.0954
Pair wise comparisons – Series 2						
Equal_Duration	Pitch-shifted -8%	2	0.252	0.244	-0.2258	0.7307
Equal_Duration	Pitch-shifted +8%	2	1.234	0.280	0.6850	1.7834
Equal_Duration	Frequency-shifted 500Hz	2	0.625	0.294	0.0492	1.2013
Equal_Duration	Contour-maintained Vocoded	2	0.344	0.286	-0.2174	0.9056
Unequal_Duration	Pitch-shifted -8%	2	0.682	0.207	0.2856	1.0885

Unequal_Duration	Pitch-shifted +8%	2	0.885	0.216	0.4613	1.3093
Unequal_Duration	Frequency-shifted 500Hz	2	0.564	0.202	0.1678	0.9606
Unequal_Duration	Contour-maintained Vocoded	2	0.727	0.213	0.3097	1.1443

Note: Lower CL and Upper CL represent the lower and upper 95% confidence limits (CL) of the confidence interval. If zero is part of that confidence interval, the treatment combination Training_Group and Stimuli is not significantly different from 0. If both confidence levels are positive, then there is a bias toward correct responses. If they are both negative, then they are biased toward incorrect responses. Each of the corrected pairwise multiple comparison test sets is separated into clusters within the table. Bold indicates significance.

Table A3 Post-hoc test results of Binomial GLM for the interaction of Time-scaled Test_Treatment & Training_Sound

Stimuli	Training_Sound	Series	Estimate	SE	Z-ratio	P-value
PC (sound A & B separated)						
Pair wise comparisons between training sounds - Series 1						
Duration stretched 50%	EqualA - EqualB	1	0.0594	0.345	0.172	0.8631
Duration stretched 50%	UnequalA - UnequalB	1	0.6801	0.355	1.915	0.1111
Duration compressed 50%	EqualA - EqualB	1	0.3645	0.350	1.043	0.3961
Duration compressed 50%	UnequalA - UnequalB	1	-0.7906	0.353	-2.238	0.1009
Pair wise comparisons between test stimuli - Series 1						
Training - Duration stretched 50%	EqualA	1	1.10e+00	0.383	2.867	0.0124
Training - Duration compressed 50%	EqualA	1	7.92e-01	0.387	2.047	0.0609
Duration compressed 50% - Duration stretched 50%	EqualA	1	3.05e-01	0.350	0.871	0.3835
Training - Duration stretched 50%	EqualB	1	1.44e+00	0.402	3.573	0.0005
Training - Duration compressed 50%	EqualB	1	1.44e+00	0.402	3.573	0.0005
Duration compressed 50% - Duration stretched 50%	EqualB	1	-4.05e+05	0.344	0.000	0.9999

Training - Duration stretched 50%	UnequalA	1	9.58e-01	0.412	2.325	0.0201
Training - Duration compressed 50%	UnequalA	1	1.88e+00	0.407	4.620	<.0001
Duration compressed 50% - Duration stretched 50%	UnequalA	1	-9.20e-01	0.357	-2.582	0.0148
Training - Duration stretched 50%	UnequalB	1	1.44e+00	0.391	3.693	0.0007
Training - Duration compressed 50%	UnequalB	1	8.93e-01	0.395	2.259	0.0358
Duration compressed 50% - Duration stretched 50%	UnequalB	1	5.50e-01	0.352	1.563	0.1179

Pair wise comparisons between training sounds - Series 2

Duration stretched 20%	EqualA - EqualB	2	4.27e-01	0.379	1.127	0.8471
Duration stretched 20%	UnequalA - UnequalB	2	-3.22e-01	0.403	-0.800	0.8471
Duration compressed 20%	EqualA - EqualB	2	-7.40e-02	0.385	-0.192	1.0000
Duration compressed 20%	UnequalA - UnequalB	2	-9.20e-06	0.416	0.000	1.0000

Pair wise comparisons between test stimuli - Series 2

Training - Duration stretched 20%	EqualA	2	4.88e-01	0.406	1.201	0.3444
Training - Duration compressed 20%	EqualA	2	5.62e-01	0.404	1.390	0.3444
Duration compressed 20% - Duration stretched 20%	EqualA	2	-7.39e-02	0.385	-0.192	0.8476
Training - Duration stretched 20%	EqualB	2	1.32e+00	0.426	3.087	0.0061
Training - Duration compressed 20%	EqualB	2	8.89e-01	0.433	2.053	0.0601
Duration compressed 20% - Duration stretched 20%	EqualB	2	4.27e-01	0.379	-1.127	0.2597
Training - Duration stretched 20%	UnequalA	2	7.05e-01	0.427	1.650	0.2967
Training - Duration compressed 20%	UnequalA	2	3.83e-01	0.440	0.870	0.4236
Duration compressed 20% - Duration stretched 20%	UnequalA	2	3.22e-01	0.403	0.800	0.4236
Training - Duration stretched 20%	UnequalB	2	6.11e-01	0.458	1.334	0.2734
Training - Duration compressed 20%	UnequalB	2	6.11e-01	0.458	1.334	0.2734
Duration compressed 20% - Duration stretched 20%	UnequalB	2	-1.27e-05	0.416	0.000	1.0000

Note: Response variables in GLMMs: the proportion of trials that birds respond with pecking A or B. Each of the corrected pairwise multiple comparison tests is separated into clusters within the table. Bold indicates significance.

Table A4 Estimates and 95% confidence intervals for the recognition of Time-scaled stimuli

Training_Group	Stimuli	Series	emmean	SE	CL (95%)	
					Lower	Upper
Log(Cor/Inco) ~ Training_Group + Stimuli + Training_Group:Stimuli + Training_Sound + Training_Sound:Stimuli + (1 BirdID), (Sound A & Sound B combined)						
Pair wise comparisons - Series 1						
Equal	Duration stretched 50%	1	0.852	0.240	0.381	1.32
Equal	Duration compressed 50%	1	0.763	0.227	0.318	1.21
Unequal	Duration stretched 50%	1	1.050	0.259	0.541	1.56
Unequal	Duration compressed 50%	1	0.629	0.229	0.181	1.08
Pair wise comparisons - Series 2						
Equal	Duration stretched 20%	2	1.21	0.277	0.668	1.75
Equal	Duration compressed 20%	2	1.41	0.286	0.849	1.97
Unequal	Duration stretched 20%	2	1.480	0.255	0.985	1.98
Unequal	Duration compressed 20%	2	1.520	0.253	1.023	2.02

Note: Lower CL and Upper CL represent the lower and upper 95% confidence limits (CL) of the confidence interval. If zero is part of that confidence interval, the treatment combination Training_Sound and Stimuli are not significantly different from 0. If both confidence levels are positive, then there is a bias toward correct responses. If they are both negative, then they are more biased toward incorrect responses. Each of the corrected pairwise multiple comparison tests is separated into clusters within the table. Bold indicates significance.

Table A5 Estimates and 95 % confidence intervals for the correct identification of time-scaled stimuli

Training_Sound	Stimuli	Series	emmean	SE	CL (95%)	
					Lower	Upper
Log(Cor/Inco) ~ Training_Group + Stimuli + Training_Group:Stimuli + Training_Sound + Training_Sound:Stimuli + (1 BirdID), (Sound A & Sound B separated)						
Pair wise comparisons - Series 1						
EqualA	Duration stretched 50%	1	1.192	0.351	0.5049	1.8794
EqualA	Duration compressed 50%	1	1.024	0.314	0.4086	1.6400
EqualB	Duration stretched 50%	1	0.511	0.288	-0.0539	1.0757
EqualB	Duration compressed 50%	1	0.501	0.288	-0.0640	1.0655
UnequalA	Duration stretched 50%	1	1.804	0.404	1.0117	2.5962
UnequalA	Duration compressed 50%	1	0.352	0.304	-0.2433	0.9474
UnequalB	Duration stretched 50%	1	0.295	0.285	-0.2627	0.8536
UnequalB	Duration compressed 50%	1	0.906	0.304	0.3105	1.5023
Pair wise comparisons - Series 2						
EqualA	Duration stretched 20%	2	1.535	0.396	0.7592	2.3108
EqualA	Duration compressed 20%	2	1.609	0.412	0.8019	2.4169
EqualB	Duration stretched 20%	2	0.886	0.345	0.2109	1.5619
EqualB	Duration compressed 20%	2	1.209	0.355	0.5127	1.9051
UnequalA	Duration stretched 20%	2	1.359	0.333	0.7059	2.0112
UnequalA	Duration compressed 20%	2	1.514	0.339	0.8483	2.1790
UnequalB	Duration stretched 20%	2	1.609	0.351	0.9199	2.2974
UnequalB	Duration compressed 20%	2	1.525	0.341	0.8578	2.1928

Note: Lower CL and Upper CL represent the lower and upper 95% confidence limits (CL) of the confidence interval. If zero is part of that confidence interval, the treatment combination Training_Sound and Stimuli are not significantly different from 0. If both confidence levels are positive, then there is a bias toward correct responses. If they are both negative, then they are more biased toward incorrect responses. Bold indicates significance.

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