

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

Ning, Z.

Citation

Ning, Z. (2024, May 2). *The flexible listener: exploring zebra finch sensitivity to spectral and temporal sound features*. Retrieved from https://hdl.handle.net/1887/3750255

Version:	Publisher's Version
License:	Licence agreement concerning inclusion of doctoral thesis in the Institutional Repository of the University of Leiden
Downloaded from:	https://hdl.handle.net/1887/3750255

Note: To cite this publication please use the final published version (if applicable).



Zebra Finches Demonstrate Cognitive Flexibility in Using Phonology and Sequence of Syllables in Auditory Discrimination

Chapter

Zhi-Yuan Ning · Henkjan Honing · Carel ten Cate

This chapter is published in Animal Cognition DOI: 10.1007/s10071-023-01763-4

ABSTRACT

Zebra finches rely mainly on syllable phonology rather than on syllable sequence when they discriminate between two songs. However, they can also learn to discriminate two strings containing the same set of syllables by their sequence. How learning about the phonological characteristics of syllables and their sequence relate to each other and to the composition of the stimuli is still an open question. We compared whether and how the zebra finches' relative sensitivity for syllable phonology and syllable sequence depends on the differences between syllable strings. Two groups of zebra finches were trained in a Go-Left/Go-Right task to discriminate either between two strings in which each string contained a unique set of song syllables ('Different-syllables group') or two strings in which both strings contained the same set of syllables, but in a different sequential order ('Same-syllables group'). We assessed to what extent the birds in the two experimental groups attend to the spectral characteristics and the sequence of the syllables by measuring the responses to test strings consisting of spectral modifications or sequence changes. Our results showed no difference in the number of trials needed to discriminate strings consisting of either different or identical sets of syllables. Both experimental groups attended to changes in spectral features in a similar way, but the group for which both training strings consisted of the same set of syllables responded more strongly to changes in sequence than the group for which the training strings consisted of different sets of syllables. This outcome suggests the presence of an additional learning process to learn about syllable sequence when learning about syllable phonology is not sufficient to discriminate two strings. Our study thus demonstrates that the relative importance of syllable phonology and sequence depends on how these features vary among stimuli. This indicates cognitive flexibility in the acoustic features that songbirds might use in their song recognition.

INTRODUCTION

Not only humans, but also songbirds learn their vocalizations early in life from their parents or other individuals. Vocal learning implies the presence of advanced auditory processing, including perception, memorization, and production of complex strings of sounds. Most emphasis in studies of vocal learning and auditory processing in birds is on the processes involved in learning the phonology, i.e., the spectro-temporal structure, of syllables, rather than on learning the syllable sequences (Vernes *et al.*, 2021).

Songbird species show a large diversity in how syllables are arranged within songs. Some songbird species, such as the canary (*Serinus canaria*) (Lehongre *et al.*, 2008), European starling (*Sturnus vulgaris*) (Eens, 1997), or willow warbler (*Phylloscopus trochilus*) (Gil & Slater, 2000) have a repertoire of syllables that are ordered in varying sequences to form phrases that together make up the song. The sequence of syllables sung within a given song is rarely an exact replicate of the previous song or of a sequence produced by the model from which the syllables are copied. This is in contrast to the vocalizations in species such as the white crowned sparrow (*Zonotrichia leucophrys*) (Soha & Marler, 2001), the chaffinch (*Fringilla coelebs*) (Riebel & Slater, 1999), song sparrow (*Melospiza melodia*) (Marler & Peters, 1987), or the zebra finch (*Taeniopygia guttata*) (Eales, 1985), in which songs consist of rather fixed sequences of syllables, and in which copied songs show limited element sequence divergence from the song models. The fact that these songbirds as well as others faithfully copy both the spectro-temporal structure of song syllables as well as the is equences, implies they have the ability to perceive and learn the phonology as well as the sequencial order of conspecific syllables in great detail.

The zebra finch is an extensively used model species for comparative studies of vocal learning as well as auditory perception. With respect to sequence learning, despite the fact that zebra finches may have certain non-learned biases as to how different syllable types are distributed over a sequence (James & Sakata, 2017), there is ample evidence that syllable sequences are affected by learning (e.g., Eales, 1985). This is supported by the finding that zebra finch songs, both in captive and wild populations, show culturally transmitted differences in the position of specific syllable types, being more similar within than between colonies (Lachlan *et al.*, 2016). Also, zebra finches first exposed to one set of syllables in a particular sequence and next exposed to a novel set, first acquire the phonological structure of the novel syllables and

next adjust the sequence of these novel syllables, indicating the involvement of at least partially different learning processes (Lipkind *et al.*, 2013). Comparable evidence of a separation between learning the phonology of syllables and learning of their sequence can also be found on other songbirds, such as the white crowned sparrows (e.g., Soha & Marler, 2001; Plamondon *et al.*, 2010).

The finding that zebra finches attend to and learn about both phonology and syllable sequence demonstrates that both are perceived and suggests that they both are relevant for communication, for instance to distinguish between individuals. However, experiments addressing which song features zebra finches use to discriminate between songs suggest a striking imbalance between the role of syllable phonology and the role of syllables sequence. For instance, Braaten et al. (2006) used an operant discrimination task (Go/Nogo) to train adult and juvenile zebra finches to discriminate the natural forward song from its reversed version (i.e., a song played backwards). Tests in which a song was presented with syllables of nonreversed phonological structure in the reversed sequence and a song in which element sequence was maintained, but the syllables were reversed, showed that the original stimuli were discriminated on the phonological structure of the syllables and not by their sequence. A recent study, also using a Go/Nogo task, investigated the role of syllable sequences versus spectro-temporal fine structure of syllables for the process of individual recognition: zebra finches were trained to discriminate songs of one male conspecific from those of four others; thereafter they were exposed to hybrid stimuli combining the syllable sequences of one individual with the spectro-temporal features of another. The results demonstrated that zebra finches mainly rely on spectro-temporal details of syllables and pay less attention to syllable sequences (Geberzahn & Derégnaucourt, 2020). A laboratory playback experiment (Mol et al., 2021) also suggested that syllable sequence is not an essential cue for recognition of familiar songs in zebra finches. In another study, Lawson et al. (2018) used a discrimination task to compare the ability of zebra finches to notice changes of syllable phonology and changes of syllable sequence in the motifs of natural songs. These results also showed that zebra finches could readily recognize the reversal of a single syllable in the motif, but largely ignore the change of syllable sequence in the motif. Similarly, zebra finches detect single syllable reversals more easily than a doubling of an inter-syllable interval (e.g., Dooling & Prior, 2017). Combined with evidence that zebra finches can detect differences between renditions of slightly different versions of the same song syllables (Fishbein et al., 2021), demonstrating the attention to fine details of the spectro-temporal structure of syllables, such findings raised the

question to what extent zebra finches attend to the sequences of syllables (Fishbein *et al.*, 2019).

Some studies have indicated that syllable sequence can play an additional role in song recognition. Lawson et al. (2018) showed that male zebra finches tested with their own songs or with those of familiar birds attended to sequences of syllables in addition to the spectrotemporal structure of these syllables. So, although zebra finches may thus show a strong bias to attend to spectro-temporal features of syllables to distinguish songs, they can also attend to syllable sequence. It suggests that more extensive experience with songs is needed before the birds acquire knowledge about syllable sequences. This was also suggested by an experiment by Braaten et al. (2006) showing that juvenile zebra finches could discriminate songs on the basis of syllable sequence alone, although this discrimination was more difficult to obtain than one based on syllable structure. However, in contrast to the studies indicating a marginal role of syllable sequences in song discrimination and suggesting that learning about sequences might be more difficult than about syllable phonology, a range of studies demonstrated that zebra finches can readily learn to distinguish strings consisting of identical syllables but differing in their sequence (e.g. van Heijningen et al., 2013; Chen & ten Cate, 2015, 2017; Chen et al., 2016; Spierings & ten Cate, 2016; Knowles et al., 2018). In a study by van Heijningen et al. (2009), zebra finches were trained in a Go/Nogo task to discriminate between stimuli in which syllables were arranged in an ABAB or an AABB sequence. They readily acquired this discrimination. When next tested with stimuli of the same sequential structures but constructed of novel exemplars of the same type of syllables (and hence differing in fine spectro-temporal details), they generalized the discrimination to the novel exemplars based on the string structure. Evidence from a neural study (Cazala et al., 2019) also using an AABB vs ABAB paradigm demonstrated that the caudomedial nidopallium (NCM) neurons encode the sequencing of syllables, which also supports the outcome of the behavioural studies described above in showing that zebra finches have no difficulty in distinguishing two strings by the sequence of their syllables. Zebra finches can thus readily use sequence information to distinguish strings differing in their sequence only.

The findings discussed above raise the question how learning about the spectro-temporal characteristics of syllables and about syllable sequences relate to each other and to the composition of the stimuli. The range of experiments mentioned above differ in methods and stimulus composition. So far, no experiment has directly compared the relative importance of

spectral structure and sequence when zebra finches have to discriminate two syllable strings that either consist of different sets of syllables or consist of the same set of syllables, but different in the sequence, using similarly structured strings and identical training and testing procedures.

In the current study we use an operant discrimination paradigm - the Go-Left/Go-Right task - to examine the relative salience of syllable phonology and syllable sequence when zebra finches must distinguish two artificially constructed 'song motifs' that are either composed of different syllable types (the 'Different-syllables group'), or two stimuli composed of the same set of syllables but differing in sequence (the 'Same-syllables group'). We investigate whether the stimulus contrast in the training affects the ease of learning by examining the speed with which the discrimination is achieved. Next, we assess to what extent the birds in the two groups attend to the syllable phonology by assessing the responses to test strings consisting of reversed syllables or of vocoded versions of these syllables. To examine the importance of syllable sequence, we assessed the responses to test strings in which the sequences are shuffled.

METHODS

Subjects

Twenty-four zebra finches (12 males, and 12 females; ages 139 - 691 days post-hatching) were used in this experiment. All birds originated 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 randomly in two experimental groups; half of the birds were assigned to the Different-syllables group, and the other half of them to the Same-syllables group (6 males and 6 females in each group; age Different-syllables group: M=309, SD=184, age Same-syllables group: M=387, SD=246). Each group was trained to discriminate between two different strings consisting of five zebra finch syllables. Within each training group one half of the birds got training strings consisting of single-element syllables, and the other half another set of stimuli consisting of one complex syllable and four single-element syllables within a string.

Operant conditioning cage

The birds were trained and tested individually in an operant conditioning cage (Skinnerbox) (70×30×45 cm) using a Go-Left/Go-Right paradigm for training and testing. A cage contained 3 pecking keys (sensors) with a red LED light at the top/bottom of each sensor (Fig. 1a). 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) 1 meter above the Skinnerbox. The volume of the speaker was adjusted to ensure that the sound amplitude in the Skinnerbox was approximately 65 dB (measured by an SPL meter, RION NL 15, RION). Sensors (S1, S2, S3), lamp, food hatch and speaker were connected to operant conditioning controller that also registered all sensor pecks.



Figure 1. (a) Schematic view of the operant conditioning apparatus (Skinner box) used for the experiment. A speaker (S) 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) is located in the upper middle of the back panel, a lamp (L) is placed at the top of the cage. Two tubes of ad libitum water (W) are placed symmetrically on two sides of the cage, three response keys (S1, S2, S3) with signal LEDs are lined horizontally in the lower middle of the back panel. (b) An example of a pair of training strings for the Different-syllables group. The birds of the Different-syllables group were trained with stimuli consisting of different syllable types: for instance, String A was the sequence of syllables A B C D E, while String B was the sequence of syllables F G H I J. (c) Modified stimuli used in the testing phase for the Different-syllables group. The birds of the training - see text for a description of these manipulations. (d) A pair of training strings for the Same-syllables group. For birds of the Same-syllables group, training stimuli consisted of the same syllables but arranged in different sequences: for instance, String A and String B consisted of the same five syllables A B C D E, but the sequences of these syllables were different between the two strings. (e) Modified stimuli in the testing phase for the Same-syllables group. These birds were also tested with 4 similarly modified versions of each training stimuli in the testing phase for the Same-syllables group. These birds were also tested with 4 similarly modified versions of each training stimuli in the testing phase for the Same-syllables group. These birds were also tested with 4 similarly modified versions of each training stimulis.

Stimuli

Training stimuli

Zebra finch syllables were selected from representative song recordings of adult males of the laboratory colony at Leiden University. The songs had not been heard before by the birds. Each string was composed of syllables belonging to different types, based on several distinctive acoustic features like the duration and spectral shape, mainly guided by the descriptions of syllable types in Lachlan *et al.* (2016). Each training string was thus consisting of five song units, each of which belonged to one of in total 13 types of single-element syllables and 8 types of complex syllables. Each bird got different combinations of syllable types as training stimuli.

The five syllables within one string were normalized in root-mean-square (RMS) amplitude and separated by a 30 ms silent interval between each two syllables to form a natural songsyllable string. The training stimuli in this experiment were 24 stimulus pairs (12 pairs for each training group), each consisting of two different strings. For the Different-syllables group, each bird was presented with a stimulus pair of which the two strings consisted of different syllable types (Fig. 1b). For the Same-syllables group, each bird was presented with a stimulus pair of which the two strings were consisting of a same set of syllables but arranged in a different sequence (Fig. 1d). To this end, we altered the syllable sequences of string A

(indicated as "A-B-C-D-E") into a different sequence "B-E-D-A-C" to construct the string B, which also avoids bigrams of syllables from string A.

When played, the strings were normalized such that the average intensity (RMS, calculated over the total duration of the stimulus) was the same for the two strings within a pair to avoid that amplitude differences affected the responses to the stimuli. The range of variation in volume recorded at the microphone was preserved. All stimuli were filtered to a bandwidth below 15kHz. All training stimuli were cut, synthesized, and filtered using Praat (version 6.1.12). The amplitude of each stimulus was adjusted by using the "Normalize" function in Audacity (version 2.3.0).

Test stimuli

To test the impact of spectral and sequential information that the birds used to discriminate the training strings, they were tested with modified versions of the training strings (Fig. 1b & 1d). We used Praat to modify each original training string to produce a version in which either the spectral features or the sequence of syllables was changed. For each training group modified stimuli were changed in an identical way (some examples of the training and test stimuli are provided as supplementary material):

- Spectrum reversal – The spectrum of each syllable in the string was reversed, but the sequence of the syllables was identical to the order in the training version. We used the "reverse selection" option in Praat to reverse the spectrum of each syllable of a training string without changing the initial order.

- Jumbled – The sequence of the syllables in the training strings of both training groups were altered from "A-B-C-D-E" to "D-C-A-E-B". For instance, if the syllable sequence of the string A in the Different-syllables group is "A-B-C-D-E", then the order manipulated version becomes "D-C-A-E-B", and the manipulated version of string B (the original sequence "F-G-H-I-J") becomes "I-H-F-J-G". Thus the "Jumbling" was applied to both string A and string B in the Different-syllables group (Fig. 1c). Likewise, this modification was applied in the Same-syllables group, by which the sequence-manipulated version of string A became "D-C-A-E-B", and the sequence manipulated version of string A became "D-C-A-E-B", and the sequence manipulated version of string A became "D-C-A-E-B", and the sequence manipulated version of string A became "D-C-A-E-B", and the sequence manipulation of string B became "A-D-B-C-E". Note that this means that the manipulated string B now has the same 1st and 5th syllables as present in training string A ("A-B-C-D-E"), since training string A and string B consisted of the same syllables.

Therefore, for the Same-syllables group, we distinguished in our analysis between the responses to "D-C-A-E-B", which will be indicated as the "Full jumbled" test string and "A-D-B-C-E" which will be indicated as "Middle jumbled" test string, and we relate the responses to these test stimuli to the responses to training string A (Fig. 1e).

- Jumbled + Spectrum reversal – This manipulation was the combination of the above Jumbled alteration and Spectrum reversal. Both the spectrum of syllables and their sequence were changed (Fig. 1b and Fig. 1d).

- Vocoded – This modification maintains the spectral (and temporal) envelope of the syllables within the string, but averages the energy within specific frequency bands, thus removing any harmonic structure. To construct these stimuli, we used the Matt Winn's Praat vocoded script (<u>http://www.mattwinn.com/praat/vocode_all_selected_v45.txt</u>) to synthesize a vocoded morph of training strings. The script was set to divide cut-off frequency bandwidths equally for 15 bands contiguous with smooth transitions (1000Hz bandwidth for one noise-vocoded band).

Procedure

We used a Go-Left/Go-Right paradigm for training and testing (Fig. 1a). The training consisted of several phases.

Acclimation and pre-training

In the acclimation phase, the birds were moved to the Skinner boxes. The food hatch remained open, so food was freely accessible in a container behind the hatch. The LED lights on the pecking sensors were on. The goal of this phase was to acclimate the bird to the cage and to show it where to find food. The bird might also already learn to peck sensors spontaneously. If in this stage the central sensor, S1, was stimulated by pecking, it would play sound string A or sound string B with a 50% chance on each. The side sensor S2 produced one of the two training strings, and the other side sensor S3 produced the other string. The LEDs of all three sensors were illuminated to attract the attention from the bird. After a few hours to one night of acclimation, the pre-training phase started by closing the food hatch. In this phase, the food hatch was closed, and the bird had to learn 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. Once the bird started to peck all the sensors regularly 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, and then to peck S2 or S3, depending on the playback sound. If the bird pecked the sensor that was linked to the stimulus being played, this was rewarded with 12 sec access to food. If the wrong sensor was pecked the light was off for 1 sec. Before any sensor was pecked, only the S1 LED was on. If the bird did not respond within 15 seconds, a trial would end automatically without food reward or light-off penalty. The duration of this phase varied from bird to bird (range: 5 - 32 days). The proportion of correct responses (see 'Analysis' section below for calculation of the 'Correct rate') was calculated on a daily basis as the individual's discrimination rate among the training stimuli.

Transition phase

When a bird learned to associate the two training sounds with the corresponding sensors and had reached a Correct rate for the training stimuli greater than 0.75 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, in which the reinforcement by food reward or darkness was reduced to occur randomly on 80% (instead of 100%) of trials. On the remaining 20% of trials, the responses were not reinforced, and the trial ended after 15 sec. If the bird kept the same level of discrimination for two days, the test phase began.

Probe testing phase

In this phase, 20% of the pecks on S1 resulted in presenting one of 10 test stimuli. These10 test stimuli were never reinforced and were randomly interspersed between training stimuli. Eight of these were modified versions of the training stimuli (four modified versions of stimulus A and four of stimulus B). The other two were non-reinforced training stimuli. The remaining 80% were training stimuli with reinforcement. Testing continued until each test stimulus had been presented 40 times to a bird. After reaching this, the bird was transferred back to its aviary. The order of stimulus presentation was random across subjects.

Analysis

For 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 two-tailed unpaired t-test (using the t-test function in GraphPad Prism 9.1.1) was used to detect differences between the two training groups.

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 a 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 'no-response' (not pecking a key). For the statistical analyses, we examined the proportion of 'correct responses' out of 'correct + incorrect responses' (Correct rate = Count_Correct / (Count_Correct + Count_Incorrect)), as well as the proportion of responses calculated as 'correct + incorrect responses' to modifications of sound A plus those to modification of sound B, as the proportion of the 40 presentations of each test stimulus (Response rate = (Count_Correct + Count_Incorrect) / (Count_Correct + Count_Incorrect) / (Count_Correct + Count_Incorrect)). In addition, we examined whether the individual test stimuli were discriminated above chance.

We used Generalized Linear Mixed-effects Models (GLMMs) to examine the discrimination of various test sounds by the birds. All model analyses were conducted in Rstudio (R Core Team, 2016). We calculated the 'Correct rate' and the 'Response rate' based on the counts of 'correct response', 'incorrect response', and 'no response', combining the response counts to (variants of) Training strings A and B, using the function cbind, R package mice; Van Buuren & Groothuis-Oudshoorn, 2011, and used these two rates as response variables in GLMMs in R (using the function glmer, R package lme4; Bates et al., 2015). We used 'Training_Group' (Same or Different syllables), 'Test_Treatment', and the interaction between these two as covariates in the full model with 'Bird_ID', 'Age', 'Number_of_Training_Trials' as the random factors and a binomial error structure of the 'Correct rate' and the 'Response rate'. The best model was chosen based on corrected Akaike criterion (AICc) provided by dredge model selection (using the function Dredge, R package MuMIn; Bartoń, 2020). The model with the smallest value of AICc was considered to be the best model by default, but if 'Training_Group' was not part of the best model, we kept it in the final model anyway because this was a variable of our interest. To determine the effect and significance of the covariates, we ran the final models and, if applicable, used Post hoc Tukey's HSD tests to make pairwise

comparisons of the test treatments (using the emmeans function, R package lsmeans; Lenth, 2016), with false discovery rate (FDR) correction of p-values (Benjamini & Hochberg, 1995) for multiple comparisons.

In the above model, the counts of the responses to (modifications of) both string A and string B were combined in all tests. This included the two test treatments 'Jumbled' and 'JumbledReversal' for both string A and B in the Same-syllables group. However, as outlined above, the jumbling of the syllables resulted in making the jumbled version of string B partly similar to training string A, and we therefore used string A as reference in this case. Because jumbling the strings for the Same-syllables group thus resulted in half of the jumbled strings being fully jumbled and the other half being middle jumbled, we also did a separate analysis for the data set of two Jumbled versions (MiddleJumbled/FullJumbled) in the Same syllables training group. In this analysis we compared the responses to training string A with those to the FullJumbled version of string are the same as those of the training string A. In this analysis, 'Test_Treatment' was used as a fixed effect in the full model to gain insight into a possible comparison among three different stimuli versions (Training/MiddleJumbled/FullJumbled). The 'Bird_ID', 'Age', and 'Number_of_Training_Trials' were included as the random factors. Here we also used a model with binomial error structure of the Correct and the Response rates.

To examine whether the birds responded above chance (50%) to each of the testing stimuli, we applied a log(correct/incorrect) as the response variables against a log (Odds-ratio) = 0 in a GLM. If correct/incorrect = 1, then the probability of observing a correct response is as large as the probability of observing an incorrect response, representing both probabilities are 0.5, then log (Odds- ratio) = log (1) = 0. Therefore, comparing the outcomes of the Binomial GLM to 0 is comparing the results to the 50% chance for a correct response.

Ethics Statement

All animal housing, care, and use was approved by the national *Centrale Commissie voor Dierproeven* (CCD) of the Netherlands and the Leiden University Animal Welfare Body (AVD number 1060020197507). None of 24 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 Skinnerboxes. During the experiment, the health

and welfare of these birds was monitored daily. The food intake of the birds was monitored daily, and additional food was given when there were signs of a low food intake.

RESULTS

Learning speed



Figure 2. Number of learning trials needed to reach the learning criterion. Individual zebra finch results are shown with open circles. There is no significant difference between the Different-syllables group and the Same-syllables group in learning speed. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range.

The discrimination training lasted until the birds reached the learning criterion of over 75% correct responses to both sound A and sound B for three successive days. All twenty-four birds finished the training and reached the learning criterion in on average 3842 (SD = 1442, N = 24) trials. No significant difference (p = 0.7733, t = 0.2916, df = 22; Fig. 2) was found between the Different-syllables group (M = 3753, SD = 1579) and the Same-syllables group (M = 3932, SD = 1283). It suggests that birds from two training groups learned approximately equally fast.

Do training groups differ in responses to test stimuli?

We compared the Correct rates and Responses rates to the training and various test stimuli between both experimental groups (Fig. 3). For the Correct rate , the best model (model 1) was chosen based on AICc (Table 1). For the Response rate, we chose the model 3 with the same factors as model 1 for the Correct rate. It was not the most recommended model by the dredge model selection, but it contained the variables of our interest and was also close to the most recommend model (AICc = 723.1, delta = 7.41, Table 1).

The only significant difference between the two training groups concerns the Correct rate for the Jumbled version (Different – Same = 0.534 ± 0.173 , p = 0.01, Table 2). There were no significant differences in the Correct rate for any of the other test stimuli between the two training groups (Fig. 3a). Note that the variation in Correct rate for the Jumbled test stimuli in the Same-syllables group is much larger than that for other test stimuli, which is caused by combining the responses to both the 'Middle Jumbled' and 'Full Jumbled' test stimuli (see below for the analysis separating among these stimuli). There were no significant differences in Response rates for any of the stimuli between two training groups (Fig. 3c).





Figure 3. Correct rate of responses and Response rate of trials a) the proportion of correct responses (Correct rate) to the training and test stimuli for the two training groups ; b) the Correct rate of responses to the training stimulus and the two Jumbled versions for the 'Same-syllables' training group; c) the Response rates to the training and test stimuli for the two training groups; d) the Response rates to the training stimulus and the two Jumbled versions for the 'Same-syllables' training group. All test stimuli got significantly lower Correct rates and significantly lower Response rates than the training stimuli. Significant differences between the responses to the various test stimuli and between the training groups are indicated: *** refers to a significant difference of $p \le 0.001$, ** refers to a significant difference of 0.001 , and * refers to a significant difference of <math>0.001 , for non-indicated comparisons <math>p value is > 0.05. Box plots show median, 1st and 3rd quartile, and whiskers the 1.5 interquartile range. The dashed line represents chance level, which was 50% for both tasks.

Table 1 Summary of the GLMs selection for (a) the proportion of correct responses if birds re	spond	to one of tw	o sounds		
and (b) the proportion of trials that birds respond with pecking A or B					
Model	df	logLik	AICc	Δi	wi
a. Correct rate of responses (sound A+B combined)					
1* Training_Group + Test_Treatment + Test_Treatment:Training_Group + (1 Bird_ID) + (1 Age) +	13	-481.009	991.5	0.00	0.964
(1 Number_of_Training_Trials)					
2 Training_Group + Test_Treatment + (1 Bird_ID) + (1 Age) + (1 Number_of_training_trials)	9	-489.805	999.2	7.79	0.020
3 Test_Treatment + (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	8	-491.141	999.6	8.13	0.017
4 Training_Group + (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	S	-789.401	1589.3	597.88	0.000
null (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	4	-790.736	1589.8	598.37	0.000
b. Response rate of trials (sound A+B combined)					
1 Test_Treatment + (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	8	-349.172	715.6	0.00	0.748
2 Training_Group + Test_Treatment + (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	9	-349.165	718.0	2.33	0.234
3* Training_Group + Test_Treatment + Test_Treatment:Training_Group + (1 Bird_ID) + (1 Age) +	13	-346.811	723.1	7.41	0.018
(1 Number_of_Training_Trials)					
4 Training_Group + (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	S	-469.717	950.0	234.32	0.000
null (1 Bird_ID) + (1 Age) + (1 Number_of_Training_Trials)	4	-469.723	947.8	232.15	0.000
Note: Best four models of the model selection (ranked by AICc and logLik) and the null models. The Akaike wei	ght (wi)	indicates the	probability	of a better	model in the
model candidates set, and Delta AICc (Δi) was used to show the difference in AICc score between the best model and choose. Only information related to both sound A and sound B were shown here, the information about the two Jum	nd the n bled ver	nodel being co sions in the 'S	mpared. A Same-syllab	* indicates bles' training	the model we group is not
displayed in this table.					

Do different test stimuli give rise to different responses?

The highest Correct and Response rates are present for the non-rewarded training stimuli. Thus, in both training groups all modifications affected the birds' responses (see Table 2). For the comparisons of responses to different test stimuli within each training group, Post hoc Tukey's HSD tests (Table 2) showed that the birds responded with a higher Correct rate and a higher Response rate to the training stimuli compared to all four testing stimuli in both training groups (Fig.3a and Fig.3c). The tests also showed that the birds of the Different-syllables training group responded with a significantly higher Correct rate to the Jumbled stimuli than to the JumbledReversal, the Vocoded stimuli and the SpectrumReversal stimuli (both p < 0.0001), and with a significantly higher Correct rate to the SpectrumReversal stimuli than to the Vocoded stimuli (p < 0.05), while the birds of the Same-syllables training group responded with a significantly lower Correct rate to the JumbledReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli than to the Jumbled stimuli (p < 0.001), the Vocoded stimuli and the SpectrumReversal stimuli (both p < 0.01).

The birds of the Different-syllables training group had lower Response rate to the Jumbled stimuli and the Vocoded stimuli than to the JumbledReversal (p < 0.01), and had a significantly higher Response rate to the SpectrumReversal stimuli than to the Jumbled stimuli and the Vocoded stimuli (both p < 0.01), while the birds of the Same-syllables training group had significantly lower Response rate to the JumbledReversal (p < 0.05), the Vocoded (p < 0.001) and the Jumbled stimuli (p < 0.01) than to the SpectrumReversal stimuli, and had a significantly higher Response rate to the JumbledReversal than to the Vocoded stimuli (p < 0.05).

To investigate the impact on discrimination of the two Jumbled versions in the Same-syllables training group, we split the data for the responses to the Jumbled version into responses to the MiddleJumbled version and FullJumbled version, comparing them with the responses given to training sound A. This showed that the birds responded with a higher Correct rate to Training sound A than to the MiddleJumbled test sound and with a higher Correct rate to the MiddleJumbled than to the FullJumbled test sound (Training – MiddleJumbled = 0.9071 ± 0.1812, MiddleJumbled – FullJumbled = 0.9094 ± 0.1603, both p < 0.001) (Fig 3b). There was no significant difference in the Response rate between these two Jumbled versions (MiddleJumbled – FullJumbled = 0.1404 ± 0.2004, p = 0.76), but both rates were lower than the Response rate to Training sound A (Training – MiddleJumbled = 1.3877 ± 0.2809, Training

- FullJumbled = 1.5281 ± 0.2783 , both p < 0.001) (Fig 3d). These results (see Table S1 in the supplementary appendix) show that the birds of the 'Same-syllables' training group pay attention to the beginning and end, as well as to the middle syllables of the strings.

Table 2 Post hoc test	results of Binomial GLMMs for th	e interaction of T	est & Train	ning_Group	
Stimuli	Training_Group	estimate	SE	z.ratio	<i>p</i> .value
a. Correct rate of responses (sound A+	B in two training groups)				
Training	Different - Same	-0.007	0.198	-0.036	0.9715
SpectrumReversal	Different - Same	0.255	0.167	1.526	0.2117
Jumbled	Different - Same	0.534	0.173	3.094	0.0100
JumbledReversal	Different - Same	0.350	0.166	2.108	0.0878
Vocoded	Different - Same	-0.020	0.168	-0.119	0.9715
Training - SpectrumReversal	Different	1.423	0.127	11.200	<.0001
Training - Jumbled	Different	0.958	0.133	7.217	<.0001
Training - JumbledReversal	Different	1.615	0.126	12.823	<.0001
Training - Vocoded	Different	1.673	0.127	13.188	<.0001
SpectrumReversal - Jumbled	Different	-0.465	0.110	-4.228	<.0001
SpectrumReversal - JumbledReversal	Different	0.192	0.102	1.884	0.0662
SpectrumReversal - Vocoded	Different	0.251	0.103	2.434	0.0186
Jumbled - JumbledReversal	Different	0.657	0.109	6.045	<.0001
Jumbled - Vocoded	Different	0.716	0.110	6.528	<.0001
JumbledReversal - Vocoded	Different	0.059	0.102	0.578	0.5631
Training - SpectrumReversal	Same	1.684	0.125	13.434	<.0001
Training - Jumbled	Same	1.499	0.127	11.781	<.0001
Training - JumbledReversal	Same	1.971	0.126	15.711	<.0001
Training - Vocoded	Same	1.660	0.127	13.076	<.0001
SpectrumReversal - Jumbled	Same	-0.185	0.102	-1.821	0.0857
SpectrumReversal - JumbledReversal	Same	0.287	0.099	2.889	0.0055
SpectrumReversal - Vocoded	Same	-0.024	0.101	-0.238	0.8117
Jumbled - JumbledReversal	Same	0.472	0.102	4.642	<.0001
Jumbled - Vocoded	Same	0.161	0.104	1.555	0.1334
JumbledReversal - Vocoded	Same	-0.311	0.101	-3.070	0.0036
b. Response rate of trials (sound A+B i	n two training groups)				
Training	Different - Same	0.118	0.467	0.252	0.9724
SpectrumReversal	Different - Same	-0.221	0.420	-0.525	0.9724
Jumbled	Different - Same	-0.185	0.412	-0.448	0.9724

JumbledReversal - Vocoded	Jumbled - Vocoded	Jumbled - JumbledReversal	SpectrumReversal - Vocoded	SpectrumReversal - JumbledReversal	SpectrumReversal - Jumbled	Training - Vocoded	Training - JumbledReversal	Training - Jumbled	Training - SpectrumReversal	JumbledReversal - Vocoded	Jumbled - Vocoded	Jumbled - JumbledReversal	SpectrumReversal - Vocoded	SpectrumReversal - JumbledReversal	SpectrumReversal - Jumbled	Training - Vocoded	Training - JumbledReversal	Training - Jumbled	Training - SpectrumReversal	Vocoded	JumbledReversal
Same	Same	Same	Same	Same	Same	Same	Same	Same	Same	Different	Different	Different	Different	Different	Different	Different	Different	Different	Different	Different - Same	Different - Same
0.3239	0.1623	-0.1616	0.642	0.319	0.480	1.756	1.432	1.594	1.114	0.466	-0.037	-0.502	0.407	-0.059	0.444	1.859	1.393	1.896	1.452	0.014	0.156
0.138	0.135	0.141	0.148	0.154	0.150	0.204	0.208	0.206	0.214	0.145	0.134	0.144	0.143	0.152	0.143	0.204	0.210	0.204	0.209	0.411	0.418
2.341	1.206	-1.142	4.353	2.071	3.191	8.621	6.879	7.746	5.197	3.217	-0.272	-3.483	2.841	-0.384	3.108	9.097	6.630	9.291	6.937	0.035	0.374
0.0275	0.2531	0.2534	<.0001	0.0480	0.0024	<.0001	<.0001	<.0001	<.0001	0.0022	0.7856	0.0010	0.0056	0.7790	0.0027	<.0001	<.0001	<.0001	<.0001	0.9724	0.9724

Note: Response variables in GLMMs: (a) the proportion of correct responses if birds respond to one of two sounds; and (b) the proportion of trials that birds respond with group weren't displayed in this table. Bold indicates significance. pecking A or B. Only information related to both sound A and sound B were shown here, the information about the two Jumbled versions in the 'Same-syllables' training

Cognitive Flexibility in Using Phonology and Sequence

Are modified stimuli still discriminated?

The above analyses concentrated on differences in the Correct rates between the groups and among the test stimuli. They don't test whether a low Correct rate also indicates that birds no longer discriminate between the modified version of training sound A and that of the similarly modified version of training sound B. If the birds are still capable of linking the modified stimuli to the respective training stimuli, the proportion of correct responses to the test stimuli should be higher than the proportion of incorrect responses. Table 3 and Fig. 4a show that for the Different-syllables group, all treatment combinations are significantly different from 0 in favour of a correct response. For the Same-syllables group, all treatments were also statistically different from 0 in favour of correct response, except the Test treatment JumbledReversal, which showed no significant difference from 0 (Fig. 4a).

For the data set of two Jumbled versions in Same-syllables group, MiddleJumbled is statistically different from 0 in favour of correct response, but FullJumbled is not significant different from 0 (Table 3), which is in line with the visualisation (Fig. 4b).



Figure 4. Visualisation of logRatios = log (Correct/Incorrect). a) For the Different-syllable group (left), all logRatios are statistically different from zero; for the Same-syllable group (right), the Test treatment JumbledReversal is not significantly different from 0; b) Results for the Jumbled test sounds of the Same-syllable group, split into Middle and Full Jumbled. For MiddleJumbled, there is a small overlap with zero; for Jumbled, it is statistically not different from 0. A * indicates that the logRatio of a Test treatment is significantly different from 0, 'ns' indicates that the logRatio 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 logRatios was based on the counts of 'correct response' and 'incorrect response' from the same data set that was also used for Figure 3.

Training Group	Stimuli	estimate	SE	CL (95)	%)
				Lower	Upper
LogRatio ~ Training_Gro	oup + Test_Treatment + Test_Treatm	nent: Training_Group	$+ (1 Bird_D) + (1 Ag$	e) + (1 Number_of_Training)	5_Trials),
data = sound A + sound E	3, n=24				
Different syllables	Training	2.090	0.142	1.812	2.368
Different syllables	SpectrumReversal	0.667	0.121	0.430	0.904
Different syllables	Jumbled	1.132	0.127	0.883	1.381
Different syllables	JumbledReversal	0.475	0.120	0.241	0.710
Different syllables	Vocoded	0.417	0.121	0.180	0.653
Same syllables	Training	2.097	0.142	1.819	2.375
Same syllables	SpectrumReversal	0.412	0.119	0.179	0.646
Same syllables	Jumbled	0.597	0.121	0.360	0.835
Same syllables	JumbledReversal	0.125	0.119	-0.108	0.359
Same syllables	Vocoded	0.436	0.121	0.200	0.673
LogRatio ~ Test_Treatm	$ent + (1 Bird_ID) + (1 Age) + (1 Nun$	aber_of_Training_Tria	als),		
data = sound A, $n=12$					
Same syllables	Training	2.073	0.309	1.468	2.678
Same syllables	MiddleJumbled	1.166	0.298	0.581	1.751
Same syllables	FullJumbled	0.257	0.293	-0.319	0.832

Table 3 Lower CL and Upper CL represent the lower and upper 95% confidence limits (CL) of the confidence interval

Note: If zero is part of the confidence interval, the treatment combination Training Group 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.

DISCUSSION

Our results show that zebra finches are capable of using both spectral features and sequential information to discriminate strings consisting of conspecific song syllables. Confirming results obtained in earlier studies on zebra finches, our study also demonstrates that zebra finches will give higher priority to using spectral features than syllable/element sequence in discrimination when the syllables differ in phonology. When strings are composed of a same set of syllables, zebra finches learn about the syllable sequence in addition to the syllable phonology.

No effect of stimulus composition on learning speed

Various studies (Braaten et al., 2006; Lawson et al., 2018; Geberzahn & Derégnaucourt, 2020) demonstrated that when zebra finches learned to discriminate between two songs, they were very sensitive to changes in the spectral domain (syllable reversals) and hardly sensitive to sequential information (sequence reversals), similar to what we observed in our 'Differentsyllables' training group. These studies indicated that the zebra finches ignored sequence cues in discrimination learning or that sequences were more difficult to learn than spectral features and might require more time. In line with this, some studies (Lawson et al., 2018; Braaten et al., 2006) indicated that if zebra finches used syllable sequences to distinguish songs, this occurred with songs to which the birds had been exposed more extensively. That learning to discriminate sequences consisting of the same sets of syllables might be more difficult than sequences consisting of different syllables was also suggested by a meta-analysis using data from 14 different acoustic Go/No-go experiments with zebra finches (Kriengwatana et al., 2016), which indicated that stimuli (either zebra finch vocalizations or human speech syllables) differing in phonetic characteristics were learned faster than those differing in sequence only. However, in our experiment, allowing a direct comparison of learning speed of comparable stimuli in identical conditions, the learning speed of the training group relying only on sequence cues is not significantly lower than that of the group trained on stimuli with different syllables. This suggests that the Same-syllables group learned about the syllable sequence in parallel with learning about the syllable phonology, without requiring more extensive exposure or training.

Cognitive flexibility in processing syllable phonology and sequence

The comparison of the correct responses to the different test stimuli showed that both training groups were similarly strongly affected by changes of the spectro-temporal features of the syllables, thus noticing such changes equally well. It demonstrates that the Same-syllables group, which can only learn a sequence of syllables when they also learn the spectro-temporal features of these syllables, gives the same weight to the spectro-temporal features as the Different-syllables group does. The difference between the two training groups concerns their responses to the jumbled test sounds. Although the jumbled test stimuli received fewer correct responses and had a lower Response rate than the training stimuli in both groups, jumbling affected the Same-syllables group much more strongly than the Different-syllables group. For the Same-syllables group, the impact of jumbling is similar to that of spectral changes. Jumbling had a lesser impact than spectral modifications in the Different-syllables group, confirming that this group mainly (although not exclusively) relied on spectral features of the syllables to distinguish the training strings. Hence, the importance of syllable sequence increased when knowledge of the sequence is needed to correctly identify different strings. This finding indicates the presence of 'cognitive flexibility' in processing string information, in which sequence learning can be added to learning of spectro-temporal features of syllables when needed to distinguish strings.

No differences were observed between the responses of both groups to reversal of the syllables and vocoding them. Reversal of syllables reverses the within-syllable spectral and amplitude pattern (i.e., any frequency changes or increasing or decreasing amplitude over an element), while vocoding maintains these patterns, but removes pitch information. Apparently, all these dimensions are taken into account for identification of syllables. Nevertheless, both groups were capable of still discriminating reversed and vocoded versions of the training stimuli, indicating that the test stimuli still maintained sufficient gross spectral differences among the syllables of a string to allow for string identification.

That full jumbling strongly affected the Same-syllables group and resulted in absence of discrimination is no surprise, as full jumbling removed all information that might relate to the original syllable sequences. However, what is of interest is that middle-jumbled also got fewer correct responses than the training stimuli, indicating that the birds were not just relying on the first and last syllables of the syllable sequence (which was suggested by studies on zebra

finches (Fishbein *et al.*, 2019) and Bengalese finches (*Lonchura striata var. domestica*) (Mizuhara & Okanoya, 2020)) but also to the sequence of the middle syllables.

Vocal production learning and discrimination learning

Altogether the results indicate that sequence learning can be 'added to' learning about spectrotemporal features of syllables if these features alone are insufficient to distinguish two syllable strings. It indicates the presence of sequence learning as a separate, but nevertheless strongly connected or partially overlapping learning process, similar to what has been observed in several studies of song production learning (Liu *et al.*, 2004; Braaten *et al.*, 2006; Lipkind *et al.*, 2013, 2017). This does not imply that song production learning and song discrimination learning rely on the same mechanisms. Song production learning occurs in male zebra finches only and only during a sensitive phase early in life, while discrimination learning can occur in both sexes and when adult. Also, vocal discrimination learning has been observed in vocal non-learning species, such as dove species (Beckers & ten Cate, 2001; Beckers *et al.*, 2003), which give attention to both spectral and temporal structure of sound strings. Hence, vocal production learning and later occurring vocal discrimination or recognition learning are likely to rely at least partly on different mechanisms.

In conclusion, our study demonstrates that although zebra finches have a bias to attend to spectral features when recognizing or discriminating strings of syllables, they can also attend to the sequence when needed. Our study did not test whether the relative importance of syllable sequence might vary if the syllable similarity between strings also varies, e.g., when not all but only part of the syllables in a string are different, or when different strings contain different exemplars of the same syllable types. It is likely that such string modifications may affect the relative weight of spectro-temporal and sequence parameters in song discrimination. Such flexibility may explain why some studies on the cues that zebra finches use to distinguish songs demonstrated absence of any impact of changes in syllable sequences on discriminating strings (Lawson *et al.*, 2018; Geberzahn & Derégnaucourt, 2020; Mol *et al.*, 2021), while other studies (van Heijningen *et al.*, 2009; Chen *et al.*, 2016; Spierings & ten Cate, 2016) showed clear sequence learning. It shows that the use of particular cues within a specific experiment should not be taken as an inability to use other cues when such cues might be useful or needed to correctly identify different strings, although the importance of the ability to also learn about syllable sequences under natural conditions remains to be elucidated. A similar flexibility, in

this case for using different spectral cues, was observed by Burgering *et al.* (2018; 2019), showing that depending on the differences among training sounds zebra finches used either pitch or spectral envelope to distinguish the training sounds. To what extend such a flexibility is also present for other song features awaits further exploration (see ten Cate & Honing, 2022). It is likely that zebra finches are not the only species that demonstrates such cognitive flexibility, although this remains to be tested. The benefit of such flexibility is that it may allow birds to adjust their perceptual tuning to those acoustic dimensions that are most relevant to distinguish songs of different individuals or other biologically relevant sounds.

Acknowledgments

We would like to thank Jeroen Hubert for help with the statistical analysis, Leda Aristidou for help with screening and synthesis of acoustic stimuli, Peter Snelderwaard for his technical assistance, and him and Michelle Geers for their animal care. Zhi-Yuan Ning was supported by the CSC (China Scholarship Council) fellowship.

References

Bartoń K. (2020). MuMIn: Multi-Model Inference. R package version 1.43.17. <u>https://CRAN.R-project.org/package=MuMIn</u>

Bates D., Maechler M., Bolker B., Walker S. (2015). Fitting linear mixed-effects models using lme4. J Stat Softw., 67(1):1-48. <u>https://doi.org/10.18637/jss.v067.i01</u>

Beckers G.J.L., ten Cate C. (2001). Perceptual relevance of species-specific differences in acoustic signal structure in Streptopelia doves. *Anim Behav.*, 62(3):511–518. <u>https://doi.org/10.1006/anbe.2001.1768</u>

Beckers G.J.L., Goossens B.M.A., ten Cate C. (2003.) Perceptual salience of acoustic differences between conspecific and allospecific vocalizations in African collared-doves. *Anim Behav.*, 65(3):605-614. <u>https://doi.org/10.1006/anbe.2003.2080</u>

Benjamini Y., Hochberg Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. J R Stat Soc Ser B., 57(1):289–300. https://doi.org/10.1111/j.2517-6161.1995.tb02031.x

Braaten R.F., Petzoldt M., Colbath A. (2006). Song Perception During the Sensitive Period of Song Learning in Zebra Finches (*Taeniopygia guttata*). J Comp Psychol., 120(2):79–88. <u>https://doi.org/10.1037/0735-7036.120.2.79</u>

Burgering M.A., ten Cate C., Vroomen J. (2018). Mechanisms underlying speech sound discrimination and categorization in humans and zebra finches. *Anim Cogn.*, 21(2):285-299. <u>https://doi.org/10.1007/s10071-018-1165-3</u>

Burgering M.A., Vroomen J., ten Cate C. (2019). Zebra finches (*Taeniopygia guttata*) can categorize vowel-like sounds both on the fundamental frequency ("Pitch") and spectral envelope. *J Comp Psychol.*, 133(1):106-117. https://doi.org/10.1037/com0000143

Cazala A., Giret N., Edeline J.M., Del Negro C. (2019). Neuronal encoding in a high-level auditory area: from sequence of elements to grammatical structure. *J Neurosci.*, 39(31):6150-6161. <u>https://doi.org/10.1523/JNEUROSCI.2767-18.2019</u>

Chen J., ten Cate C. (2015). Zebra finches can use positional and transitional cues to distinguish vocal element strings. *Behav Processes.*, 117:29-34. <u>https://doi.org/10.1016/j.beproc.2014.09.004</u>

Chen J., Jansen N., ten Cate C. (2016). Zebra finches are able to learn affixation-like patterns. *Anim Cogn.*, 19(1):65-73. <u>https://doi.org/10.1007/s10071-015-0913-x</u>

Chen J., ten Cate C. (2017). Bridging the gap: Learning of acoustic nonadjacent dependencies by a songbird. J Exp Psychol Anim Learn Cogn., 43(3):295–302. https://doi.org/10.1037/xan0000145

Derégnaucourt S., Poirier C., Kant A.V., Linden A.V., Gahr M. (2013). Comparisons of different methods to train a young zebra finch (*Taeniopygia guttata*) to learn a song. J Physiol Paris., 107(3):210–218. <u>https://doi.org/10.1016/j.jphysparis.2012.08.003</u>

Dooling R.J., Prior N.H. (2017). Do we hear what birds hear in birdsong? *Anim Behav.*, 124:283–289. <u>https://doi.org/10.1016/j.anbehav.2016.10.012</u>

Eales L.A. (1985). Song learning in zebra finches: Some effects of song model availability on what is learnt and when. *Anim Behav.*, 33(4):1293–1300. <u>https://doi.org/10.1016/S0003-3472(85)80189-5</u>

Eens M. (1997). Understanding the complex song of the European starling: an integrated ethological approach. Adv Study Anim Behav., 26:355–434. <u>https://doi.org/10.1016/S0065-3454(08)60384-8</u>

Fishbein A.R., Idsardi W.J., Ball G.F., Dooling R.J. (2019). Sound sequences in birdsong: how much do birds really care? *Philos Trans R Soc Lond B Biol Sci.*, 375(1789):20190044. https://doi.org/10.1098/rstb.2019.0044

Fishbein A.R., Prior N.H., Brown J.A., Ball G.F., Dooling R.J. (2021). Discrimination of natural acoustic variation in vocal signals. *Sci Rep.*, 11:916. <u>https://doi.org/10.1038/s41598-020-79641-z</u>

Geberzahn N., Derégnaucourt S. (2020). Individual vocal recognition in zebra finches relies on song syllable structure rather than song syllable order. *J Exp Biol.*, 223(9):jeb220087. <u>https://doi.org/10.1242/jeb.220087</u>

Gil D., Slater P.J.B. (2000). Song organisation and singing patterns of the willow warbler,
Phylloscopustrochilus.Behaviour,137(6):759-782.https://doi.org/10.1163/156853900502330

James L.S., Sakata J.T. (2017). Learning biases underlie "universals" in avian vocal sequencing. *Curr Biol.*, 27(23):3676-3682.e4. <u>https://doi.org/10.1016/j.cub.2017.10.019</u>

Knowles J.M., Doupe A.J., Brainard M.S. (2018). Zebra finches are sensitive to combinations of temporally distributed features in a model of word recognition. J Acoust Soc Am., 144(2):872. <u>https://doi.org/10.1121/1.5050910</u>

Kriengwatana B., Spierings M.J., ten Cate C. (2016). Auditory discrimination learning in zebra finches: Effects of sex, early life conditions and stimulus characteristics. *Anim Behav.*, 116:99–112. <u>https://doi.org/10.1016/j.anbehav.2016.03.028</u>

Lachlan R.F., van Heijningen C.A., Ter Haar S.M., ten Cate C. (2016). Zebra finch song phonology and syntactical structure across populations and continents—a computational comparison. *Front Psychol.*, 7:980. <u>https://doi.org/10.3389/fpsyg.2016.00980</u>

Lawson S.L., Fishbein A.R., Prior N.H., Ball G.F., Dooling R.J. (2018). Relative salience of syllable structure and syllable order in zebra finch song. *Anim Cogn.*, 21(4):467-480. <u>https://doi.org/10.1007/s10071-018-1182-2</u>

Lehongre K., Aubin T., Robin S., Del Negro C. (2008). Individual signature in canary songs: Contribution of multiple levels of song structure. *Ethology*, 114(5):425-435. https://doi.org/10.1111/j.1439-0310.2008.01486.x

Lenth R.V. (2016). Least-Squares Means: The R Package lsmeans. J Stat Softw., 69(1):1-33. <u>https://doi.org/10.18637/jss.v069.i01</u>

Lipkind D., Marcus G.F., Bemis D.K., Sasahara K., Jacoby N., Takahasi M., Suzuki K., Feher O., Ravbar P., Okanoya K., Tchernichovski O. (2013). Stepwise acquisition of vocal combinatorial capacity in songbirds and human infants. *Nature*, 498(7452):104–108. https://doi.org/10.1038/nature12173

Lipkind D., Zai A.T., Hanuschkin A., Marcus G.F., Tchernichovski O., Hahnloser R.H.R. (2017). Songbirds work around computational complexity by learning song vocabulary independently of sequence. *Nat Commun.*, 8(1):1247. <u>https://doi.org/10.1038/s41467-017-01436-0</u>

Liu W.C., Gardner T.J., Nottebohm F. (2004). Juvenile zebra finches can use multiple strategies to learn the same song. *Proc Natl Acad Sci.*, 101(52):18177–18182. <u>https://doi.org/10.1073/pnas.0408065101</u>

Marler P., Peters S. (1987). A sensitive period for song acquisition in the song sparrow, *Melospiza melodia*: A case of age-limited learning. *Ethology*, 76(2):89–100. https://doi.org/10.1111/j.1439-0310.1987.tb00675.x

Mizuhara T., Okanoya K. (2020). Do songbirds hear songs syllable by syllable? *Behav Processes.*, 174:104089. <u>https://doi.org/10.1016/j.beproc.2020.104089</u>

Mol C., Bolhuis J.J., Moorman S. (2021). Vocal learning in songbirds: the role of syllable order in song recognition. *Philos Trans R Soc Lond B Biol Sci.*, 376(1836):20200248. https://doi.org/10.1098/rstb.2020.0248

Plamondon S.L., Rose G.J., Goller F. (2010). Roles of syntax information in directing song development in white-crowned sparrows (*Zonotrichia leucophrys*). J Comp Psychol., 124(2):117-32. <u>https://doi.org/10.1037/a0017229</u>

Riebel K., Slater P.J.B. (1999). Song type switching in the chaffinch, *Fringilla coelebs*: timing or counting? *Anim Behav.*, 57(3):655-661. <u>https://doi.org/10.1006/anbe.1998.0984</u>

Soha J.A., Marler P. (2001). Vocal syntax development in the white-crowned sparrow (Zonotrichia leucophrys). J Comp Psychol., 115(2):172-80. <u>https://doi.org/10.1037/0735-7036.115.2.172</u>

Spierings M.J., ten Cate C. (2016). Budgerigars and zebra finches differ in how they generalize in an artificial grammar learning experiment. *Proc Natl Acad Sci.*, 113(27):E3977-84. <u>https://doi.org/10.1073/pnas.1600483113</u>

van Buuren S., Groothuis-Oudshoorn K. (2011). Mice: Multivariate Imputation by Chained Equations in R. J Stat Softw., 45(3):1-67. <u>https://doi.org/10.18637/jss.v045.i03</u>

van Heijningen C.A., de Visser J., Zuidema W., ten Cate C. (2009). Simple rules can explain discrimination of putative recursive syntactic structures by a songbird species. *Proc Natl Acad Sci.*, 106(48):20538–20543. <u>https://doi.org/10.1073/pnas.0908113106</u>

van Heijningen C.A., Chen J., van Laatum I., van der Hulst B., ten Cate C. (2013). Rule learning by zebra finches in an artificial grammar learning task: which rule? *Anim Cogn.*, 16(2):165-175. <u>https://doi.org/10.1007/s10071-012-0559-x</u>

Vernes S.C., Kriengwatana B.P., Beeck V.C., Fischer J., Tyack P.L., ten Cate C., Janik V.M. (2021). The multi-dimensional nature of vocal learning. *Philos Trans R Soc Lond B Biol Sci.*, 376(1836):20200236. <u>https://doi.org/10.1098/rstb.2020.0236</u>