TweetyNet: A neural network that enables high-throughput, automated annotation of birdsong

Yarden Cohen^{1@*}, David Nicholson^{2@}, Alexa Sanchioni¹, Emily K. Mallaber¹, Viktoriya Skidanova¹, Timothy J. Gardner^{3‡}

1 Biology department, Boston University, Boston, MA, USA

2 Biology department, Emory University, Atlanta, GA, USA

3 Phil and Penny Knight Campus for Accelerating Scientific Impact, University of Oregon, Eugene, OR, USA

These authors contributed equally to this work.

* vcohen1@mgh.harvard.edu timg@uoregon.edu

Abstract

Songbirds have long been studied as a model system of sensory-motor learning. Many analyses of birdsong require time-consuming manual annotation of the individual elements of song, known as syllables or notes. Here we describe the first automated algorithm for birdsong annotation that is applicable to complex song such as canary song. We developed a neural network architecture, "TweetyNet", that is trained with a small amount of hand-labeled data using supervised learning methods. We first show TweetyNet achieves significantly lower error on Bengalese finch song than a similar method, using less training data, and maintains low error rates across days. Applied to canary song, TweetyNet achieves fully automated annotation of canary song, accurately capturing the complex statistical structure previously discovered in a manually annotated dataset. We conclude that TweetyNet will make it possible to ask a wide range of new questions focused on complex songs where manual annotation was impractical.

Introduction

Songbirds provide an excellent model system for investigating sensorimotor learning [1]. Like many motor skills, birdsong consists of highly stereotyped gestures executed in a sequence [2]. In this and many other ways, birdsong resembles speech: song is learned by juveniles from a tutor, like babies learning to talk [3]. A key advantage of songbirds as a model system for studying vocal learning is that birds sing spontaneously, often producing hundreds or thousands of song bouts a day. This provides a detailed readout of how song is acquired during development, and how this skilled behavior is maintained in adulthood. Leveraging the amount of data that songbirds produce requires methods for high-throughput automated analyses. For example, automated methods for measuring 10 similarity of juvenile and tutor song across development [4,5] led to important advances 11 in understanding the behavioral [6,7] and genetic [8] bases of how vocalizations are 12 learned. These examples demonstrate how automated methods that enable analysis of 13 large-scale behavioral datasets contribute to realizing the potential of songbirds as a 14 model system. 15

However, this potential to address central questions of sensorimotor learning is 16 currently hindered by a lack of high-throughput automated methods for scaling up other 17

1

2

q

> types of analyses. The central issue is that many analyses require researchers to annotate 18 song. Annotation is a time-consuming process done by hand (typically with GUI-based 19 applications, e.g., Praat, Audacity, Chipper [9–11]). An example of Bengalese finch 20 song annotated with a GUI is shown in Fig. 1. Researchers annotate song by dividing 21 it up into segments (red lines in Fig. 1), often referred to as syllables or notes, and 22 assigning labels to those segments (letters in Fig. 1). Annotation makes several types 23 of analyses possible. For example, annotation is required to build statistical models of 24 syntax [12-15], to fit computational models of motor learning that precisely quantify how 25 single syllables change over the course of an experiment [16, 17], and to relate behavior 26 to neural activity [18-20]. Annotating song greatly increases our ability to leverage 27 songbirds as a model system when answering questions about how the brain produces 28 syntax observed in sequenced motor skills, and how the brain learns to adaptively control muscles. 30

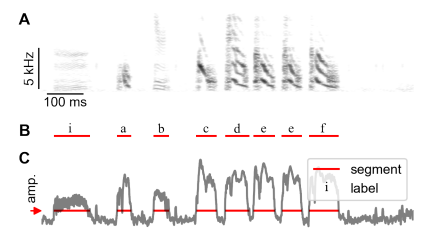


Fig 1. Annotation of birdsong. A. Spectrogram showing a brief clip of Bengalese finch song with different syllable types. **B.** Text labels over red segments are applied by human annotators to assign those segments to various syllable classes. **C.** Segments were extracted from song by finding continuous periods above a fixed amplitude threshold. Red arrow to left of panel **C** indicates the user-defined amplitude threshold.

Previous work has been done on automating annotation, as we briefly review below in 31 Proposed Method and Related Work, but these methods are challenged by the variable 32 song of some species. To illustrate these challenges, Fig 2A-C presents examples of 33 annotated songs from different species. When a species' song consists of just a few 34 syllables sung repeatedly in a fixed motif, methods based on template matching or other 35 algorithms (see Proposed Method and Related Work below) can be applied. This is 36 true for zebra finches, as can be seen in a song from one individual shown in Fig 2A. 37 However, many species have songs that are more complex than the stereotyped motif of zebra finches. Complex songs can contain a large vocabulary of syllable types arranged 30 in multiple motifs or phrases, with phrases sequenced according to complex transition statistics. For example, Bengalese finch song contains "branch points", where a given 41 syllable may transition to more than one other class of syllable. An example of a branch 42 point is indicated above the spectrogram in Fig 2B. In addition, Bengalese finch song 43 can contain syllables that repeat, with the number of repeats varying from rendition to 44 rendition. Both branch points and repeats prevent existing algorithms from effectively 45 annotating Bengalese finch song (Fig 2E). Canary song is even more complex (Fig 2C). Some individuals may have as many as 50 unique classes of syllables in their repertoire. 47 Bouts of canary song can last more than a minute instead of a few seconds (Fig 2D).

These long songs contain individual syllable types that can be very short, under 10ms, or 49 very long, ranging up to 500ms (Fig 2F). Some syllables are very quiet, and others loud. 50 Because of this extreme range of amplitude, common methods for segmenting audio of 51 song into syllables can fail. Segments are typically defined as points where the smoothed 52 sound envelope or other song-related acoustic features [4] stay above some threshold, 53 indicated by the dashed lines in Fig 3. In the case of canary song, if sound energy or 54 other acoustic features are filtered on timescales short enough to accurately segment the 55 shortest syllables, then the longest syllables will be subdivided. This problem is also 56 commonly encountered when analyzing the variable songs of young zebra finches. Fig 3 57 illustrates how canary song is difficult to segment in an automated manner. Finally, canary song has a hierarchical structure where syllables occur in trilled repetitions, called phrases, that themselves obey long-range syntax rules [12, 21]. Phrases can differ in duration depending on the type of syllable being repeated and similarly inter-syllable 61 silent gaps vary widely in duration (Fig. S1). Because of all this complexity, there are 62 currently no automated methods for accurate annotation of canary song. 63

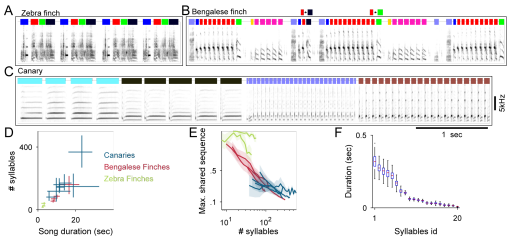


Fig 2. The challenge of annotating complex songs. A. The zebra finch repeating motif allows annotation by matching its template spectrogram without segmenting different syllables (colored bars). B. Bengalese finch songs segmented to syllables shows variable transitions and changing numbers of syllable repeats. C. A third of one domestic canary song of median duration segmented to syllables reveals repetitions (phrase) structure. D. The median, 0.25 and 0.75 quantiles of song durations (x-axis) and of syllables per song (y-axis) for 2 canary strains, Bengalese finches and Zebra finches (color coded) E. Variable songs are not suited for template matching. Songs contain repeating sequences of syllables but because of sequence variability songs with more syllables (x-axis) share smaller sequence fractions (y-axis) F. Distributions of syllable duration for one domestic canary. The bird had 20 different syllable types (x-axis, ordered by mean syllable duration). Box plot shows median, 0.25 and 0.75 quantiles of syllable durations. Whiskers show the entire range.

Proposed Method and Related Work

Previous work has been done to automate annotation, as referenced above, that we now briefly review. The crucial point here is that none of the methods work for canary song, for the reasons we outlined and demonstrated in Figs. 2 and 3, necessitating the development of an algorithm like the one we present. However, for birdsong that consists largely of a single fixed motif, like that of zebra finches, several methods have

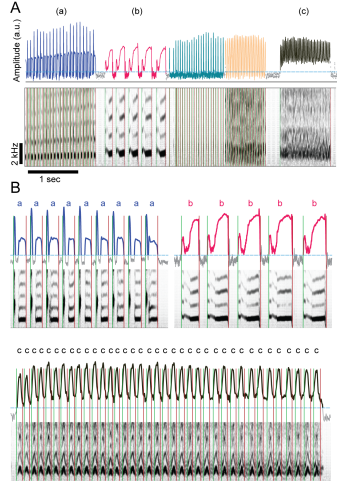


Fig 3. Examples of failure to segment canary song. A. Several seconds of domestic canary song, presented as a spectrogram, beneath a plot of a band-pass filtered sound amplitude. To segment song, an amplitude threshold can be taken, marked by the dashed line on the amplitude trace, and then an automated program finds continuous segments of above-threshold amplitude and marks the onset and offset times of those segments (green, red lines in the spectrogram panel). B. Focusing on three examples (a-c matching panel A), segmenting by threshold crossing with a fixed filtering bandwidth does not work well for canaries. Above threshold amplitudes are shown in bold colored lines and reveal that syllables of type 'a' are broken into 2 components and syllables of type 'c' are not separated by low amplitude.

been widely used, including semi-automatic clustering methods [22,23], and template 70 matching [24–26]. Several studies have also applied supervised learning algorithms to 71 annotation, such as Hidden Markov Models [27], k-Nearest Neighbors [28], and support 72 vector machines [29]. These algorithms can annotate more variable song with branch 73 points and repeats, like that of Bengalese finches, but they all require segmenting song 74 to extract the engineered features used to train the algorithms (e.g. acoustic parameters 75 like pitch and duration). To our knowledge there has been no large-scale comparison of performance of these different algorithms, but at least one study suggests they may not 77 generalize well across songs of different individuals [30]. Additionally, feature extraction can fail if segmentation is noisy, e.g. because of changes in audio equipment set-up, 79 background noises, etc. Here again we stress that canary song exhibits wide ranges 80 in amplitude, and often requires annotators to set multiple thresholds to successfully 81 extract segments. These factors contribute to the lack of automated algorithms for 82 annotating canary song. 83

Given these issues, we sought to develop an algorithm for automated annotation 84 that (1) can learn features from data, and (2) does not require segmented syllables 85 to predict annotations. To meet both these criteria, we developed an artificial neural 86 network that we call TweetyNet, shown in (Fig 4). TweetyNet takes as input windows 87 from spectrograms of song and produces labels for each time bin of that spectrogram 88 window. TweetyNet requires no pre-processing of song spectrograms - most importantly, 89 segmentation of song into syllables is not needed. Silent gaps between syllables are 90 labelled in training data, and these silent labels are assigned to gaps between syllables 91 when TweetyNet inference is applied to a new song. 92

Essentially, the network combines two types of layers found in neural networks:(1) convolutional layers, common in computer vision tasks to learn features of images [31-33], and (2) recurrent layers, often used to predict sequences [34]. A recurrent layer is a natural choice because the input image or spectrogram is defined by two axes (time and frequency) with very different correlation structure. Specifically, the temporal dimension of songbird vocalization, like music and environmental noises, contains regularities in multiple time scales that are unrelated to the regularities of the frequency axes. The bidirectional LSTM (Long-Short-Time-Memory) recurrent layer is designed to capture 100 these temporal correlations. [35, 36]. 101

To predict annotation, we feed consecutive windows from spectrograms to trained 102 networks and then concatenate the output vectors of labeled timebins. Finally, we simply 103 find uninterrupted runs of a single syllable label to annotate song syllables from this 104 framewise classification. As discussed below, this final step can include a "debounce" 105 step that requires a minimum syllable duration and choosing a single label for consecutive 106 time bins not labeled as silence by majority vote. In the rest of the results below we 107 show that this simple method trained end-to-end provides robust predictions of segment 108 onsets, offsets, and labels. 109

Surprisingly, beyond the work previously cited, we find little research that addresses 110 the problem of learning to classify each time bin of a vocalization, either for human 111 speech or birdsong. The architecture we present here is somewhat similar to early deep 112 networks models for speech recognition, but a crucial difference is that state-of-the-art 113 models in that area map directly from sequences of acoustic features to sequences of 114 words [37]. The success of these state-of-the-art models is attributed to the fact that they 115 learn this mapping from speech to text, **avoiding** the intermediate step of classifying 116 each frame of audio, as has previously been shown [34]. In other words, they avoid the 117 problem of classifying every frame that we set out to solve. The architecture that we 118 develop is most directly related to those that have been used for event detection in audio 119 and video [35, 36] and for phoneme classification and sequence labeling [34, 38]. The 120 closest prior model for segmenting and labeling birdsong is [39]. Several aspects of that 121 study provide context for the contributions of our work. The authors compared different 122 pipelines that combine a neural network for recognizing syllable segments with Hidden 123 Markov Models that learns to predict syllable sequences, and in this way improve the 124 output of the network. They measured performance of these pipelines on a large dataset 125 of hand-annotated Bengalese finch song which they made publicly available [40]. 126

In summary, the key prior art is the important work of Koumura and Okanoya [39]. 127 This work anticipates the overall structure of our model, but through the integration of 128 multiple distinct components that are individually optimized. In contrast, TweetyNet 129 is a single neural network trained end-to-end, meaning it does not require optimizing 130 multiple models. Below we show that TweetyNet meets our criteria for an algorithm 131

93

94

95

96

97

98

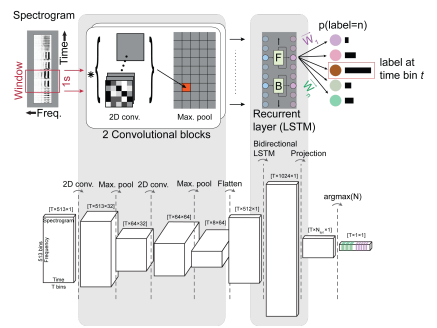


Fig 4. TweetyNet neural network architecture. Top, network schematic. TweetyNet takes as input a window, specified in time bins, from a spectrogram (red box, left) and in a sequence of steps (left to right) outputs a label for each time bin within the window: (1) The convolutional blocks produce a set of feature maps by performing a cross-correlation-like operation (asterisk) between their input and a set of learned filters (greyscale boxes). A max-pooling operation down samples the feature maps. (2) The recurrent layer is made up of Long Short Term Memory (LSTM) units, and the number of units equals the number of time bins in the spectrogram window. This step is designed to capture dependencies across time using both forward (F) and backward (B) passes through time to learn. (3) A projection $(\vec{W}_{t,s})$ onto the different syllable classes, s = 1..n, resulting in a vector of probabilities at each time bin t that the label is n. The number of classes, n, is predetermined by the user and includes a class for no-song time bins. (4) Each time bins is labeled by choosing the class with the highest probability and the labelled time bins are used to separate continuous song segments from no-song segments and to annotate each song-segment with a single label. Bottom, the shapes of tensors (multi-dimensional arrays) that result from each operation the network performs.

that learns features from data and does not require segmented song to make predictions. 132 To do so we we benchmark TweetyNet on Bengalese finch and canary song, and where 133 possible compare the performance to [39]. Additionally we show that we achieve robust 134 performance: across songs of individuals, which can vary widely even within a species; 135 across many bouts of song from one individual, e.g. across days of song, and; across 136 multiple species. Lastly we show that this performance required only a small amount of 137 manually annotated data to train TweetyNet models accurately enough to recreate and 138 add details to the deep structure of canary syntax. 139

Results

140

TweetyNet annotates Bengalese finch song with low error rates across individuals.

We first set out to test whether our network robustly annotates syllables across a large 143 number of individual birds. To do so, we made use of the publicly available repository 144 of Bengalese Finch song [40], used to benchmark hybrid neural network-HMM models 145 from [39] as referenced in Proposed Method and Related Work. The repository contains 146 song from 10 individual birds, with hundreds of bouts of hand-annotated song for each 147 bird. Each individual's song had different number of syllables and obeyed a different 148 syntax. To benchmark TweetyNet models on this dataset, we generated learning curves 149 that plot error of the model as a function of the size of the training set (duration in 150 seconds). The learning curves give us an estimate of the smallest amount of hand-labeled 151 training data we would need to obtain the lowest error that the TweetyNet model can 152 achieve. For each bird we split the data into fixed training and test sets, with durations 153 of 900 and 400 seconds respectively. Then for each training set duration we trained 154 10 replicates with a randomly-drawn subset of the training data. We computed error 155 metrics for each training replicate on the held-out test set for each individual. (See 156 Materials and methods for details.) As shown in Fig 5, these learning curves demonstrate 157 that TweetyNet models achieved low error rates across all ten birds. We first looked at 158 frame error, a percentage that measures the number of times the label predicted by the 159 model for each time bin in a spectrogram did not match the ground truth label. For all 160 birds TweetyNet models achieve less than 8% frame error with the smallest training set 161 duration of 30 seconds (Fig 5A). From the learning curve we can estimate that across 162 birds, the lowest frame error that TweetyNet models produce is roughly 4%, and that 163 they achieve this with just 180 seconds (three minutes) of training data. (For specific 164 values, see Table 1.) Larger training sets did not further reduce error. 165

To better understand how well the network segments and labels songs, we used 166 another metric, the syllable error rate, which is analogous to the word error rate that is 167 widely used in the speech recognition literature. This metric is an edit distance that 168 counts the number of edits (insertions and deletions) needed to convert a predicted 169 sequence of syllables into the ground-truth sequence. The error rate is normalized by 170 dividing it by the length of the sequences for comparison across birds (e.g. if one bird 171 sang more syllables per bout than another). Measuring the syllable error rate confirmed 172 that TweetyNet consistently achieved similar error rates across the ten birds, as shown in 173 Fig 5B. Because this metric was also used in [39] (as "note error rate"), we can compare 174 our results directly to theirs. As indicated by blue circles in Fig 5B, the best-performing 175 models in that study achieved syllable error rates of 0.83 and 0.46 with two and eight 176 minutes of training data, respectively. TweetyNet always achieved much lower syllable 177 error rates. Taken together, the results from benchmarking TweetyNet on this dataset 178 indicate that the architecture performs well across the song of many individual birds. In 179 addition, it dramatically outperforms existing models with less training data, and does 180 so while being trained end-to-end without requiring optimizations of multiple steps in a 181 pipeline. 182

TweetyNet models achieve low error across days even when trained with just the first three minutes of song recorded.

We next sought to benchmark TweetyNet in a scenario similar to long-term behavioral experiments for which we hope to automate annotation. For this purpose we used another publicly-available repository [41] with hand-labeled song from four Bengalese finches. Importantly, the repository contains most or all of the songs sung by each 188

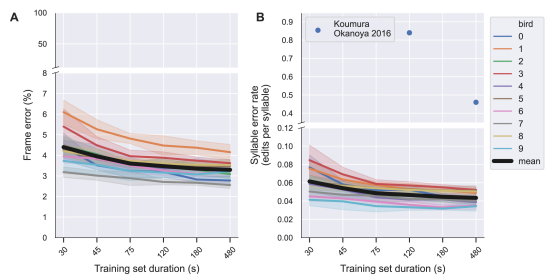


Fig 5. TweetyNet annotates song with low error rates across ten individal Bengalese finches. Model 'learning curves' showing the reduction in annotation error (y-axis) on a held-out test set as a function of the size of the training set (x-axis). Shown are the 'frame error rate' (A) measuring the percent of mislabeled time bins and the 'syllable error rate' (B) measuring the normalized sequence edit distance. Each colored line corresponds to one bird from dataset. The solid line indicates mean error across ten training replicates for each training set duration, and the translucent error band around the solid lines indicates standard deviation. Thicker black lines indicate the mean across birds. Circular blue markers indicate mean syllable error rate across birds reported in [39] for a different algorithm using the same dataset.

bird for multiple consecutive days, as is typically done during a long-term behavioral 189 experiment, and annotation for all those songs (recall that experimenters usually are able 190 to annotate only a limited number). Here we sought to measure how well TweetyNet 191 models would perform when an experimenter takes the *first* set of songs of some duration 192 n and annotates those songs manually before using them to train a network. This 193 stands in contrast to the experiment in Fig. 5, where we trained multiple replicates with 194 random subsets of songs from a larger training set, in order to obtain a better estimate 195 of expected error rates. Of course our goal is to avoid the need for experimenters to 196 label a large dataset by hand and then use it to train multiple replicates with random 197 subsets of that data, just to find the best performing network. If we show that we can 198 achieve comparable error rates with just the first n minutes of song, we can be more 199 confident that TweetyNet models will robustly segment and label hours of song recorded 200 across days. 201

Using the learning curves in Fig 5 we estimated that three minutes of data was the 202 shortest duration training set we could use to obtain the lowest error rate achieved by 203 models. Thus, we trained single TweetyNet models with the first three minutes of song 204 sung by a bird on one day, and then measured the accuracy of that model using all 205 other songs across multiple days. The test datasets we used to obtain these measures 206 were in almost all cases at least as large as those we used to benchmark models in the 207 learning curves. The mean duration of these test datasets was 1528 seconds (standard 208 deviation of 888.6 seconds, i.e. 25 minutes mean, 14 minutes standard deviation), in 209 contrast to Fig 5 where we measured error with a test set of 400 seconds (6 minutes 210 40 seconds). Hence this approach gave us multiple estimates of how a single trained 211 model performs on relatively large datasets. TweetyNet models trained in this manner 212 did achieve low frame error (Fig 6A) and low syllable error rates (Fig 6B) across days without exhibiting large fluctuations. The frame error ranged from 2-4% across 3-5 days of song, comparable to those observed when training with a random subset of songs, as in Fig 5. In one case, for one bird, the frame error did increase on the last day, but was still within the low end of the range seen for all birds, and this increase did not appear to translate into an increase in the syllable error rate (Fig 6B and Fig 6C, bird ID or60yw70, red line).

We also found that TweetyNet models trained on the first three minutes of song 220 maintained a low syllable error rate across days (Fig 6B and Fig 6C), again comparable 221 to what we observed in the learning curves (Fig 5B). Here we additionally tested whether 222 a simple post-processing step could further lower the error rate. This "majority vote" 223 transform consists of taking each labeled segment (bordered by two segments that the 224 network predicted were "unlabeled" / "silent" segments), finding the label occurred 225 most frequently within that segment, and then assigning that label to all time bins 226 within the segment. As shown in Fig 6C, this simple post-processing step did lower 227 the syllable error rate of TweetyNet models. We did not find that this post-processing 228 step had a large effect on the frame error (not shown in plot), from which we infer 229 that this transform removes small frame errors (e.g. a single time bin) that give rise 230 to spurious extra segments, and correcting these in turn produces a large drop in the 231 syllable error rate. Hence we have shown using Bengalese finch song that TweetyNet 232 outperforms existing models and that, with only minimal cleaning of its output, analyses 233 of behavioral experiments can be scaled up to very large datasets. 234

TweetyNet annotates minutes-long canary songs with low error ²³⁵ rates across individuals ²³⁶

After demonstrating TweetyNet's high performance across multiple individuals of the same species and across multiple songs of individual birds, we wanted to test TweetyNet across species. We chose the domestic canary (serinus canaria) - a species for which there are no published annotation algorithms and whose rich song repertoire offers a unique opportunity for neuroscience research [12, 21, 42–44].

As in our first test in Bengalese finches, we curated training sets of 1-10 minutes of song from 3 canaries and measured the frame error rates in a held-out test set 20-30 minutes long. (Training sets are relatively longer than the Bengalese tests since canary songs can be up to a minute or more in length and even sparse sampling of the full repertoire requires these longer training sets.) Still, Fig 7 shows that in three canaries the model learning curves asymptote with 8-10 minute training sets to frame error rates similar to TweetyNet's performance in Bengalese finches.

Unlike TweetyNet's performance in Bengalese finches, the frame error rates in annotating canary songs cannot be compared to alternative algorithms using published data and results. Furthermore, the length of these songs, usually containing hundreds of syllables, mean that even in very low error rates we expect annotation errors in many songs (Table 1, Fig. S2). These annotation errors can occur at the onset of song and in transitions between canary phrases (Fig 8) and affect analyses of canary syntax. 249

To gauge the effect of such errors, in the next section we evaluate the accuracy of 255 syntax models estimated from TweetyNet's automatic annotation. 256

Automated analysis of canary song structure.

Sequences of canary phrases contain transitions with different 'memory' depths. Namely, the probability distribution of transition outcomes from a given phrase is captured by Markov chains with variable lengths. As shown in a recent study in Waterslager canaries, this syntax structure is captured parsimoniously by probabilistic suffix trees 260 261 262

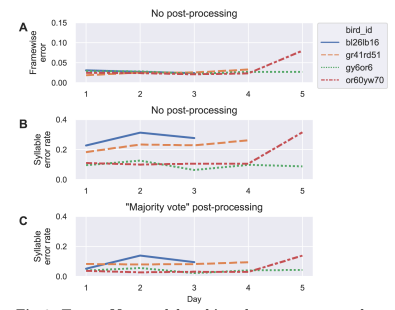


Fig 6. TweetyNet models achieve low error across days of Bengalese finch song, even when trained with just the first three minutes of song recorded. A. TweetyNet models trained on the first three minutes of song from day 1 achieved low frame error across days. The mean duration of the set of songs for each day that we used to measure error was 1528 seconds(888.6 S.D.), (i.e. 25 minutes (14 minutes S.D.)), Different line colors and styles indicate individual birds **B**. TweetyNet models trained on the first three minutes of song from day 1 also demonstrate a low syllable error rate across days. **C**. The syllable error rates in **B** further improve after applying a "majority vote" post-processing (assigning the dominant label in each continuous segment of time bins not annotated as 'silence', see methods). For one bird (or60yw70), the error did increase on the last day, but was still within the low end of the range seen for all birds.

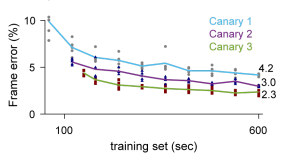


Fig 7. TweetyNet segments and labels canary song with low error rates, similar to Bengalese finches, across individuals. Models were trained on 60s-600s of song from each individual. The mean frame error (lines) of five models (markers) trained with different randomly-drawn subsets from the training set was measured on a separate 1500-2000s test set from each individual. The asymptotic error rates, annotated to the right of the curves, overlaps with the error rates in the Bengalese finch data sets

(PST) [12, 45]. The root node in these graphical models, appearing in the middle of Fig 9A,B and Fig 10A,B, represents the zero-order Markov, or base rate, frequencies of the different phrases, labelled in different colors and letters. Each branch, emanating 264

dataset	training set duration (s)	frame error (%)	syllable er- ror rate	syllable error rate (majority vote)	% Near Boundary
B.F. 1	120	$3.5 {\pm} 0.5$	$0.05 {\pm} 0.01$	n/a	n/a
B.F. 1	180	$3.4{\pm}0.5$	$0.04{\pm}0.01$	n/a	n/a
B.F. 1	480	$3.3{\pm}0.5$	$0.04{\pm}0.01$	n/a	n/a
B.F. 2	180	$2.9{\pm}1.4$	$0.2{\pm}0.09$	$0.06 {\pm} 0.04$	64.9 ± 14.3
Can.	240	$3.9{\pm}1.0$	$0.155 {\pm} 0.072$	$0.076 {\pm} 0.037$	51.1 ± 10.7
Can.	600	$3.1{\pm}0.8$	$0.09 {\pm} 0.016$	$0.051 {\pm} 0.011$	58.3 ± 11.3
Can.	6000	$2.1{\pm}0.8$	$0.069 {\pm} 0.013$	$0.031 {\pm} 0.005$	68.6 ± 13.9

Table 1. Error metrics of TweetyNet models for different species and training set sizes For each Bengalese finch (BF) and canary (Can) data set we evaluate test-set errors metrics for models trained on several training-sets sizes (measured in seconds). Presented are the mean \pm standard deviation across all birds and experiment replicates. The *frame error rate* and *syllable error rate* columns present the raw error shown in learning curves (Figs. 5,7). The *syllable error rate (majority vote)* column shows the syllable error rate after applying post-hoc cleaning of annotation, where we assigned a single label to each segment by majority vote and discarded all segments below a set duration (methods). The *% Near Boundary* column shows the percent of frame errors involving silent periods that occur within 0-2 time bins of syllable boundaries (onsets and offsets, see Materials and methods).

from the colored letters in Figs 9,10 represents the set of Markov chains that end in the specific phrase type designated by that label. For example, the 'A' branch in Fig 9a includes the first order Markov model 'A' and the second order Markov chains 'FA' and '1A' representing the second order dependence of the transition from phrase 'A'. These models are built by iterative addition of nodes up the branch to represent longer Markov chains, or a transition's dependence on longer sequences of song history.

Figures 9 and 10 demonstrate that TweetyNet parses domestic canary song with 271 an accuracy sufficient to extract its long-range order. In both of these figures, we set 272 parameters of the PST estimation algorithm to derive the deepest syntax structure 273 possible without overfitting as practiced in a recent study [12] that used about 600 hand 274 annotated songs of Waterslager canaries. In this example, using 2.2% of the data set, 275 about 40 songs, to train a TweetyNet model and predict the rest of the data reveals the 276 deep structures shown in Fig 9B - comparable to using 600 hand annotated songs of the 277 same bird (Fig 9A). With more training data, Tweetynet's accuracy improves as does 278 the statistical strength of the syntax model. In Fig 10B a TweetyNet model was trained 279 on 19% of the data, about 340 songs, and predicted the rest of the data. The resulting 280 syntax model can be elaborated to greater depth without overfitting. To crosscheck 281 this deeper model, we manually annotated all 1764 songs of that bird, revealing a very 282 similar syntax model (Fig 10A). 283

In sum, we find that TweetyNet, trained on a small sample of canary song, is accurate enough to automatically derive the deep structure that has formed the basis of recent studies [12, 46].

Larger data sets of annotated canary song add details and limit the memory of the syntax structure 288

The increase in syntax detail, presented in Fig 10, is possible because more rare nodes can be added to the PST without over-fitting the data. Formally, the PST precision 290

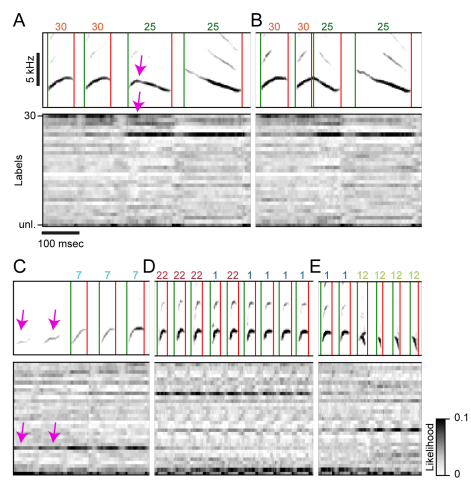


Fig 8. Variants of canary song introduce segmentation and annotation errors. Canary vocalizations contain variations that challenge TweetyNet. The examples in panels A-E show spectrograms on top of the time-aligned likelihood (gray scale) assigned by a trained TweetyNet model to each of the labels (y-axis, 30 syllable types and the tag *unl*. for the unlabeled segments). Green and red vertical lines and numbers on top of the spectrograms mark the onset, offset, and labels predicted by the model. **A,B**. Transitions between syllables can occur without a silence gap. In this example, TweetyNet assigns higher likelihood to both syllables (c.f. pink arrow). In rare variants the model ignores the first syllable (A) **C**. Syllables produced weakly or deformed still get higher likelihood. **D**. Transition between phrases of very similar syllables ($22 \rightarrow 1$) introduce label confusion. **E**. Canaries can produce completely overlapping syllables. The model assigns high likelihood to both classes but is forced to choose only one

increase in larger data sets is defined by the decrease in minimal node frequency allowed in the process of building PST models (Fig 11), as measured in model cross validation (methods). In our data set, we find an almost linear relation between the number of songs and this measure of precision - close to a tenfold precision improvement. 291 292 293 294

In Fig 10A, this increased precision allowed reliably adding longer branches to the PST to represent longer Markov chains (in comparison to Fig 9A). In this example, using a dataset 3 times larger revealed a 5-deep branch that initiate with the beginning of song 297

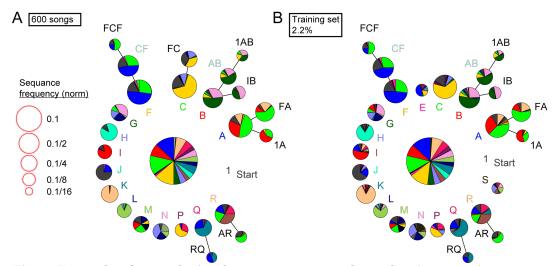


Fig 9. Example of reproducing long-range syntax dependencies, seen in *Waterslager* canaries, in another strain using a TweetyNet model trained on a small fraction of the data. A. Long-range order found in 600 domestic canary songs annotated with human proof reader (methods, similar dataset size to [12]). Letters and colors label different phrase types. Each branch terminating in a given phrase type indicates the extent to which song history impacts transition probabilities following that phrase. Each node corresponds to a phrase sequence, annotated in its title, and shows a pie chart representing the outgoing transition probabilities from that sequence. The nodes are scaled according to their frequency (legend). Nodes that can be grouped together (chunked as a sequence) without significantly reducing the power of the model are labeled with blue text. **B.** The songs used to create the PST in A are a subset of 1764 songs. A TweetyNet model was trained using about 2.2% of that dataset (about 9.5% of the data in A). The PST created from the model's predicted annotation of the entire dataset is very similar to A.

('1ABGN') indicating a potential global time-in-song dependency of that transition. The PST in Fig 10A also has branches that did not 'grow' when more songs were analyzed (e.g. the 'B', 'Q', and 'R' branches) - indicating a potential cutoff of memory depth that is crucial in studying the neural mechanisms of song sequence generation.

The data sets used in Figs 9A,10A, and Fig 11, are about 10 times larger than previous studies. To ascertain the accuracy of the syntax models, in creating the data sets we manually proof read TweetyNet's results (see methods). Across 5 different human proof readers we compare the time required to manually annotate canary song with the proof reading time and find that using TweetyNet saves 95-97.5 percent of the labor.

Taken together, the TweetyNet algorithm allowed us to annotate many more songs of individual complex singers than previously demonstrated, with high accuracy across individuals and across species. This accuracy allowed fully-automated analyses, saved most of the labor, and revealed novel details of canary syntax in a new strain.

Discussion

The family of songbirds that learns by imitation consists of over 4500 species of birds. ³¹³ Some of these singers, such as the canary, produce songs that are much too complex ³¹⁴ to be automatically annotated with existing methods, and for these complex singers ³¹⁵

307

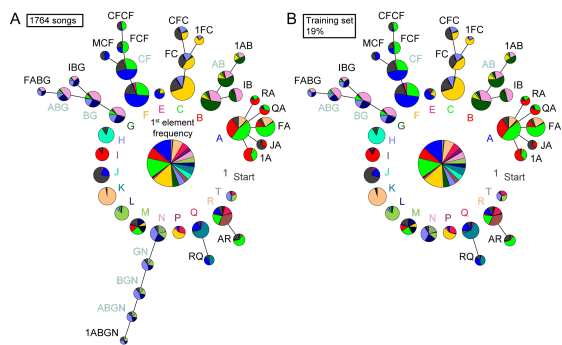


Fig 10. Example of how using TweetyNet to process a larger dataset of canary song adds detail and limits the memory of the syntax structure. A. The full dataset of 1764 songs from Fig 9, annotated with a human proof reader, allowed creating a PST with greater detail. Compared to Fig 9A, some branches did not grow. B. An almost identical PST was created *without* a human proof reader from a TweetyNet model trained on 19% of the data. The fluctuation in transition probabilities accumulates in long sequences and, in this example, increased the minimal sequence probability included in the PST. This difference prevented the inclusion of the 'N' branch.

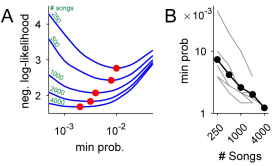


Fig 11. Using datasets more than 5 times larger than previously explored increases statistical power and the precision of syntax models. A. Ten-fold cross validation is used in selection of the minimal node probability for the PSTs (x-axis). Lines show the mean negative log-likelihood of test set data estimated by PSTs in 10 repetitions (methods). Curves are calculated for datasets that are sub sampled from about 5000 songs. Red dots show minimal values - the optimum for building the PSTs. **B.** The decrease in optimal minimal node probability (y-axis, red dots in panel A) for increasing dataset sizes (x-axis) is plotted in gray lines for 6 birds. The average across animals is shown in black dots and line.

little is known about the syntax structure and organization of song. Even for birds with simple adult songs, a detailed description of song development will require the application of new methods. This is particularly true for early song development where template based extraction of song syllables and clustering of syllable forms provides an incomplete picture of the full variability of song.

A recent study illustrated the surprises that await a more detailed analysis of song. 321 The canary, one of the most widely bred species of domesticated songbird, was recorded 322 for 2 hours or more and hundreds of songs were manually annotated and cross validated. 323 This data set revealed a new complexity to the statistical structure of canary song - the 324 song follows long-range rules where specific subsets of syllables follow transition statistics 325 governed by 4th and 5th order Markov processes in phrase types [12]. This rich behavior 326 motivated another recent study to implant miniature microscopes in singing canaries, 327 and the recorded neural signals included hierarchical memory traces corresponding to the 328 complex syntax [46]. The sophistication of the neural representation of song in canaries 329 was largely unanticipated based on decades of neural recordings in simpler singers. 330

The present project was motivated by these recent studies and the knowledge that new fundamental discoveries in vocal learning and neural dynamics will follow if automated annotation of complex song becomes possible. Some methods for automated annotation exist, but previous work suggests these methods have their own limitations, especially when applied to song with many syllable types and variable sequence such as that of Bengalese finches and canaries.

The TweetyNet algorithm described here is a work in progress, with many clear 337 paths for improvement. Still the first syllable error rates described here are dramatic 338 improvements over a prior model for song parsing. We used publicly-available datasets 339 of Bengalese finch song to benchmark TweetyNet. We showed that it achieves low error 340 rates across many individuals. On Bengalese finch data, our single network trained end-341 to-end performs better than a previously proposed hybrid HMM-neural network model 342 and does so with less training data (Fig. 5). We then showed that TweetyNet models 343 achieve low error across days, in thousands of Bengalese finch songs, even when trained 344 with just the first three minutes of song. This experiment, while strongly restricting the 345 data available for model training, demonstrates the usefulness of TweetyNet in 'real-life' 346 laboratory settings - for experimentalists that want to hand annotate as little as possible. 347

We next reported that TweetyNet was sufficiently accurate to reproduce the recent 348 findings on the complex syntax structure of canary song with fully automated machine-349 classified song. Specifically, a TweetyNet model trained on just 10 minutes of canary song 350 could accurately recover the statistical structure reported from 600 manually annotated 351 songs - exceeding 100 minutes. Furthermore, a deep network trained on 340 annotated 352 songs, about 19% of the data, could classify a larger data set of more than 1700 songs 353 and build a much more complete statistical model of song revealing additional depth to 354 the long-range syntax rules and extending prior reports on the complexity of the canary 355 song behavior. This more complex statistical model was validated using a manually 356 curated data set of all songs. 357

With a trained model performing at this level it becomes feasible to examine the 358 effect of social context on song syntax, circadian variations in syntax, or the effects of 359 distinct neural perturbations that could effect song syntax while keeping syllable forms 360 intact. On top of sequence variations, many song studies require syllable similarity 361 metrics to examine the effects of such neural or song pertubations, or the ontogeny 362 of syllable forms through development. Here we used TweetyNet to classify the most 363 likely syllable in every time point, focusing not on variations in syllable form but the 364 sequential structure of song syntax. But, the syllable classification is the final processing 365 step in TweetyNet achieved by maximum a-posteriori (MAP, or argmax) estimation 366 following the calculation of similarity to all possible syllables. Thus, the full likelihood 367 function that TweetyNet produces prior to classification may itself be a useful metric for syllable structure, allowing for example the time course of syllable form to be examined through development or as a result of neural perturbations. A syllable similarity metric that can be assigned at each point in time or frame of a spectrogram without syllable segmentation is, by itself, a new development in the field and can be used, in future development, to improve TweetyNet and to apply it to many more species whose song is difficult to segment.

To make TweetyNet useful to a large research community, we developed the vak 375 library - a user-friendly toolbox that enables researchers to apply TweetyNet simply by 376 adapting existing configuration files. This library does not require extensive programming 377 knowledge or expertise in neural networks. The framework will allow users to explore 378 different methods of optimizing neural network models that might improve segmentation, 379 and also generate alternative architectures that could incorporate distinct features and 380 topologies. For example, in many domains transformer networks have recently replaced 381 LSTMs for sequence processing. Substituting transformer layers for the LSTM layer could 382 provide advances here. [47]. Aspects of other deep networks applied to animal motor 383 control may improve TweetyNet. Examples include object detection architectures [48,49] 384 applied to mouse ultrasonic vocalizations and animal motion tracking, and generative 385 architectures applied to birdsong and other vocalizations [50-52]. Lastly we note that in 386 principle TweetyNet and vak library can be applied to any other annotated vocalization, 387 including calls of bats, mouse ultrasonic vocalizations, and dolphin communication. We 388 do not claim to have achieved the best possible method for automated annotation of 389 vocalizations with neural networks using supervised learning methods, although we have 390 aimed to establish a strong baseline for the work that will build upon ours. That said, we 391 are confident our method enables songbird researchers to automate annotation required 392 for analyses that address central questions of sensorimotor learning. 393

Materials and methods

Ethics declaration

All procedures were approved by the Institutional Animal Care and Use Committees of Boston University (protocol numbers 14-028 and 14-029). Song data were collected from n = 5 adult male canaries. Canaries were individually housed for the entire duration of the experiment and kept on a light–dark cycle matching the daylight cycle in Boston (42.3601 N). The birds were not used in any other experiments.

Data availability

Open datasets of annotated Bengalese finch song are available at <https://figshare. 402 com/articles/BirdsongRecognition/3470165> and <https://figshare.com/articles/ Bengalese_Finch_song_repository/4805749>. Audio data sets of canary song are 404 available from the corresponding author on request. 405

Code availability

The code implementing the TweetyNet architecture, and code to reproduce figures in this paper, are available at <https://github.com/yardencsGitHub/tweetynet> (version 0.4.3, 10.5281/zenodo.3978389). To aid with reproducibility of our experiments, and to make TweetyNet more accessible to researchers studying birdsong and other animal vocalizations, we developed a software library, vak, available at <https://github.com/ 411 NickleDave/vak>. Both TweetyNet and vak are implemented using the following open- source scientific Python libraries: torch [53], torchvision [54], numpy [55,56], scipy [57],

395

394

406

> dask [58], pandas [59], matplotlib [60, 61], seaborn [62], jupyter [63], attrs [64] and 414 tqdm [65]. 415

Data collection

Use of available datasets

Bengalese finch song is from two publicly-available repositories. The first [40] was used for 418 results in 4 and can be found at <https://figshare.com/articles/BirdsongRecognition/ 3470165>. It accompanied the paper [39]. The second [41] was used for results in 420 Fig 5 can be found at <https://figshare.com/articles/Bengalese_Finch_song_ 421 repository/4805749>. Apart from recordings made for this manuscript we used pub-422 licly available datasets of Waterslager canary songs [12], Bengalese finch songs [39] and 423 Zebra finch songs [66]. 424

Domestic canary song screening

Birds were individually housed in soundproof boxes and recorded for 3-5 days (Audio-426 Technica AT831B Lavalier Condenser Microphone, M-Audio Octane amplifiers, HDSPe 427 RayDAT sound card and VOS games' Boom Recorder software on a Mac Pro desktop 428 computer). In-house software was used to detect and save only sound segments that 429 contained vocalizations. These recordings were used to select subjects that are copious 430 singers (≥ 50 songs per day) and produce at least 10 different syllable types. 431

Domestic canary audio recording

All data used in this manuscript was acquired between late April and early May 2018 – a 433 period during which canaries perform their mating season songs. Birds were individually 434 housed in soundproof boxes and recorded for 7-10 days (Audio-Technica AT831B Lavalier 435 Condenser Microphone, M-Audio M-track amplifiers, and VOS games' Boom Recorder 436 software on a Mac Pro desktop computer). In-house software was used to detect and 437 save only sound segments that contained vocalizations. Separate songs were defined by 438 silence gaps exceeding 1 second.

Audio processing

Segmenting annotated phrases of Waterslager canaries

The dataset of waterslager canaries was available from a previous project in the Gardner 442 lab [12]. These songs were previously segmented into phrases, trilled repetitions of 443 syllables, and not to individual syllables. To include this data in Fig 2 we needed to break annotated phrase segments into syllable segments. In each segmented phrase, we 445 separated vocalization and noise fluctuations between vocalizations by fitting a 2-state 446 hidden Markov model with Gaussian emission functions to the acoustic signal. The 447 suspected syllable segments resulting from this procedure were proofread and manually 448 corrected using a GUI developed in-house (https://github.com/yardencsGitHub/ 449 BirdSongBout/tree/master/helpers/GUI). 450

Preparing data sets of domestic canaries

Bootstrapping annotation with TweetyNet In this manuscript we used annotated 452 domestic canary datasets an order of magnitude larger than previously published. To 453 create these datasets we used TweetyNet followed by manual proofreading of its results. 454 This process, described below, allowed 'bootstrapping' TweetyNet's performance. 455

416

417

432

425

444

451

440

Song syllables were segmented and annotated in a semi-automatic process:

- A set of 100 songs was manually segmented and annotated using a GUI developed in-house (https://github.com/yardencsGitHub/BirdSongBout/tree/master/ helpers/GUI). This set was chosen to include all potential syllable types as well as cage noises. 460
- The manually labeled set was used to train TweetyNet (https://github.com/ yardencsGitHub/tweetynet). 461
- In both the training phase of TweetyNet and the prediction phase for new annotations, data is fed to TweetyNet in segments of 1 second and TweetyNet's output is the most likely label for each 2.7msec time bin in the recording.
- The trained algorithm annotated the rest of the data and its results were manually verified and corrected. 467

Assuring the identity and separation of syllable classes The manual steps 468 in the pipeline described above can still miss rare syllable types of mislabel sylla-469 bles into the wrong classes. To make sure that the syllable classes are well sepa-470 rated all the spectrograms of every instance of every syllable, as segmented in the 471 previous section, were zero-padded to the same duration. An outlier detection al-472 gorithm (IsolationForest: <https://scikit-learn.org/stable/modules/generated/ 473 sklearn.ensemble.IsolationForest.html>) was used to flag and re-check potential 474 mislabeled syllables or previously unidentified syllable classes. 475

Preparing spectrograms inputs for TweetyNet Spectrograms were created from 476 audio files using custom Numpy (Bengalese finch) or Matlab (canary) code. All spectro-477 grams for song from a given species were created with the same parameters (e.g., number 478 of samples in the window for the Fast Fourier Transform). From initial studies we found 479 that it was necessary to perform standard transforms on spectrograms such as a log 480 transform in order for the neural network to learn. We did not notice any difference in 481 the nature of the transform (i.e., we also used $\log + 1$) although here we do not study 482 this systematically. 483

Network Architecture

The network takes a 2D window from a spectrogram as input (red box, left in Fig 4) 485 and produces as output labels for each time bin in the window. The spectrogram 486 window passes through two standard convolutional blocks, each of which consists of 487 a convolutional layer and a max pooling layer. The convolutional layer performs a 488 cross-correlation like operation (asterisk in Fig 4) between the spectrogram window and 489 learned filters (greyscale boxes in Fig 4) to produce feature maps. The max pooling 490 layer uses a similar operation to further reduce feature maps to maximum values within 491 a sliding window (orange bin in Fig 4). Importantly, the window size we use in the max 492 pooling layer has a "width" of one time bin, so that this layer does not down-sample 493 along the time axis (although the convolutional layer does). The output of the second 494 convolutional block passes through a recurrent layer made up of LSTM units, where the 495 number of units equals the number of time bins in the spectrogram window.

The final layer in TweetyNet is a projection $(\overline{W}_{t,s})$ of the recurrent layer's output onto the different syllable classes, s = 1..n, resulting in a vector of n syllable-similarity scores for each spectrogram time bin t. The number of classes, n, is predetermined by the user and includes a class for no-song time bins. At present this non-song class includes both background noises and silence, and future iterations of the model may solution to the different syllable classes, s = 1..n, resulting in a vector of n syllable-similarity scores for each spectrogram time bin t. The number of classes, n, is predetermined includes a class for no-song time bins. At present this non-song class solution the different syllable classes and silence for the model may solution to the different syllable classes includes by the user and includes a class for no-song time bins. At present this non-song class includes both background noises and silence, and future iterations of the model may

484

> distinguish between these for better performance. To segment syllables, the bin-wise 502 syllable-similarity scores are first used to select a single syllable class per time bin by 503 choosing the label with the highest syllable-similarity score. Since similarity scores can 504 be normalized, this is akin to maximum a-posteriori (MAP) label selection. Then, the 505 labelled time bins are used to separate continuous song segments from no-song segments 506 and to annotate each song-segment with a single label using majority decision. 507

Training and benchmarking TweetyNet

Benchmarking of TweetyNet was performed with the vak library. We apply standard 509 methods for benchmarking supervised machine learning algorithms, following best 510 practices [67]. We leverage functionality of the vak library that extends best practices 511 for benchmarking to the domain where where dataset size is measured in duration, as 512 described in Learning curves. 513

Data transformations

As stated above, the input to the network consists of spectrogram windows. To produce 515 this input, we slid a window of fixed length across spectrograms, essentially creating 516 an array of every possible window from each spectrogram. This array was randomly 517 permuted then fed to the network in minibatches during training, along with the expected 518 output, vectors of labels for each timebin in the spectrogram windows. These vectors 519 of labeled timebins are produced programmatically by vak from annotations consisting 520 of segment labels and their onset and offset times. For Bengalese finch song we used 521 windows of 88 time bins, and for canary song we used windows of 370 time bins. We 522 carried out preliminary experiments where we varied the window size for Bengalese finch 523 song, but did not find that larger windows greatly increased accuracy, although they did 524 increase training time. 525

Learning curves

For the studies shown in Figs. 5,7, we created learning curves, that display a metric such 527 as frame error rate as a function of the amount of training data. For each individual 528 bird we fit networks with training sets of increasing size (duration in seconds) and then 529 measured performance on a separate test set. 530

In the case of Bengalese finches, we used training sets with durations ranging from 531 30-480 seconds. For each network trained, audio files were drawn at random from a 532 fixed-size total training set of 900 seconds until the target size (e.g. 60 seconds) was 533 reached. If the total duration of the randomly drawn audio files extended beyond the 534 target duration, they were clipped at the target duration in a way that ensured all 535 syllable classes were still present in the training set. For each bird we trained ten 536 replicates, where each replicate had a different subset of randomly-drawn audio files to 537 create the target training set size. For all Bengalese finches, we measured accuracy on a 538 separate test set with a fixed size of 400s. We chose to use a totally-separate fixed-size 539 set (instead of e.g. using the remainder of the training data set) so we could be sure 540 that any variance in our measures across training replicates could be attributed to the 541 randomly-drawn training set, and not to changes in the test set. We computed metrics 542 such as frame error rate and syllable error rate on the held-out test set for each bird. 543

For canaries we used test set duration of 1500-2000 seconds and training sets of 544 60-600 seconds for the learning curves in Fig. 7. For the result in Table 1 we used a 545 test set of 5000 seconds and a training set of 6000 seconds. The method for generating 546 learning curves as just described is built into the vak library and can be reproduced 547

514

508

using its *learncurve* functionality in combination with the configuration files we shared 548 (reference link) and the publicly-available datasets. 549

Metrics

We measured performance with two metrics. The first is the frame error rate, that simply 551 measures for each acoustic frame (in our case each time bin in a spectrogram) whether 552 the predicted label matches the ground truth label. Hence the range of the frame error 553 rate is between 0 and 1, i.e. can be stated as a percent, and gives an intuitive measure of 554 a model's overall performance. Previous work on supervised sequence labeling, including 555 bidirectional-LSTM architectures similar to ours, has used this metric [34,38]. 556

The second metric we used is commonly called the word error rate in the speech 557 recognition literature, and here we call it the syllable error rate. This metric is an edit 558 distance that counts the number of edits (insertions and deletions) needed to convert 559 a predicted sequence into the ground-truth sequence. The error rate is normalized by 560 dividing it by the length of the sequences. 561

In Table 1 we provide two additional measures. The first is a lower bound on the 562 percent of all frame errors that can be attributed to slightly-misaligned syllable onsets 563 and offsets. These syllable boundaries are naturally variable in creating the ground truth 564 hand annotated data sets. Spectrogram time bins in which a trained TweetyNet model 565 and the ground truth disagree and only one of them assigns the 'unlabeled' tag can 566 potentially be around segment boundaries. In Fig. S2 we show the histogram of distances, 567 in spectrogram bins, of such frame errors from ground truth segment boundaries. The 568 majority is concentrated in 0-2 bins away from the boundaries, amounting the overall 569 percents summarized in Table 1. The second is syllable error rate after applying post-hoc 570 cleaning of annotation. This cleanup is done in two steps: (1) discard all segments 571 shorter than 5msec (using 10 msec adds an insignificant improvement in some birds) and 572 (2)assign a single label to each segment of time bins not labeled as 'silence' by majority 573 vote. 574

Model output as syllable likelihoods

In Fig 8 we present model outputs one step prior to assigning the most likely label to each 576 spectrogram time bin. At that stage, one before the argmax(N) step in Fig 4, the model 577 output for a given time bin t is a real-valued affinity $a(t,s) \in \mathcal{R}$ of all predefined syllable 578 classes s. In Fig 8 we convert these numbers to likelihoods by subtracting the minimum 579 value and normalizing separately for each time bin $L(t,s) = \frac{a(t,s) - \min_{s'} a(t,s')}{\sum_{\sigma} [a(t,\sigma) - \min_{s'} a(t,s')]}$. This transformation was done for each time bin $L(t,s) = \frac{a(t,s) - \min_{s'} a(t,s')}{\sum_{\sigma} [a(t,\sigma) - \min_{s'} a(t,s')]}$. 580 transformation was done for presentation only. Applying the commonly-used softmax 581 transform $(x \to \frac{exp(x)}{\sum_x exp(x)})$ is equivalent since we only keep the maximal value. 582

Data analysis - song structure

Shared template dependence on number of syllables in song (Fig 2e)

In each bird we define an upper bound for repeating parts of songs using pairwise 585 comparisons. For each song we examined all other songs with equal or larger number of 586 syllables and found the largest shared string of consecutive syllables. The fraction of shared syllables is the ratio between the number of shared sequence and the number of 588 syllables in the first, shorter, song. Then, we bin songs by syllable counts (bin size is 10 589 syllables) and calculate the mean and standard deviation across all pairwise comparisons. 590

550

587

583

584

Probabilistic suffix tree (Figs 9,10)

For each canary phrase type we describe the dependency of the following transition 592 on previous phrases with a probabilistic suffix tree. This method was described in a 593 previous publication from our lab (Markowitz et. al. 2013, code in https://github. 594 com/jmarkow/pst). Briefly, the tree is a directed graph in which each phrase type is a 595 root node representing the first order (Markov) transition probabilities to downstream 596 phrases, including the end of song. The pie chart in Figs 9.10 shows such probabilities. 597 Upstream nodes represent higher order Markov chains that are added sequentially if 598 they significantly add information about the transition. 599

Model cross validation to determine minimal node frequency

To prevent overfitting, nodes in the probabilistic suffix trees are added only if they 601 appear more often than a threshold frequency, P_{min} . To determine P_{min} we replicate the 602 procedure in [12] and carry a 10-fold model cross validation procedure. In this procedure 603 the dataset is randomly divided into a training set, containing 90 percent of songs, and 604 a test set, containing 10 percent of songs. A PST is created using the training set and 605 used to calculate the negative log likelihood of the test set. This procedure is repeated 606 10 times for each value of P_{min} , the x-axis in Fig 11a. For data sets of different sizes 607 (curves in Fig 11a, x-axis in Fig 11b) the mean negative log-likelihood across the 10 608 cross validation subsets and across 10 data sets, y-axis in Fig 11a, is then used to find 609 the optimal value of P_{min} - the minimum negative log-likelihood that corresponds to the 610 highest precision without over-fitting the training set. All PSTs in Figs 9,10 are created 611 using the cross-validated P_{min} . 612

Acknowledgments

This study was supported by NIH grants R01NS104925, R24NS098536 (T.J.G.) We thank J. Markowitz and T.M. Otchy for sharing song datasets, and Nvidia Corporation for a technology grant (Y.C., Sober lab).

Supporting information

Fig. S1 Consecutive canary phrases can include acoustically-similar syllables but differ in the duration of inter-syllabic gaps.

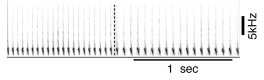


Fig S1. Example of two consecutive canary phrases that differ mostly in inter-syllable gaps. In this case, annotation methods that first segment syllables and then use acoustic parameters to classify them will introduce errors. By simultaneously learning acoustic and sequence properties, TweetyNet overcomes this weakness.

Fig. S2 Most errors of trained TweetyNet models are disagreement on 520 syllable boundaries of 0-2 time bins. 621

613

591

600

617

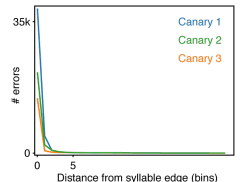


Fig S2. Potential syllable boundary disagreements are time bins in which the ground truth test set or the trained TweetyNet model disagree and just one of them assigns the 'unlabeled' silence tag. The histograms show the distances of those time bins from the

nearest syllable boundary in test sets 5000 second long.

References

- 1. Mooney R. Neurobiology of song learning. Current opinion in neurobiology. 2009;19(6):654–660.
- Fee MS, Scharff C. The songbird as a model for the generation and learning of complex sequential behaviors. ILAR journal. 2010;51(4):362–377.
- Brainard MS, Doupe AJ. What songbirds teach us about learning. Nature. 2002;417(6886):351–358.
- Tchernichovski O, Nottebohm F, Ho CE, Pesaran B, Mitra PP. A procedure for an automated measurement of song similarity. Animal behaviour. 2000;59(6):1167– 1176.
- Mets DG, Brainard MS. An automated approach to the quantitation of vocalizations and vocal learning in the songbird. PLoS computational biology. 2018;14(8):e1006437.
- Tchernichovski O, Mitra PP, Lints T, Nottebohm F. Dynamics of the vocal imitation process: how a zebra finch learns its song. Science. 2001;291(5513):2564– 2569.
- 7. Mets DG, Brainard MS. Learning is enhanced by tailoring instruction to individual genetic differences. eLife. 2019;8.
- 8. Mets DG, Brainard MS. Genetic variation interacts with experience to determine interindividual differences in learned song. Proceedings of the National Academy of Sciences. 2018;115(2):421–426.
- 9. Praat: doing Phonetics by Computer;. Available from: https://www.fon.hum. uva.nl/praat/.
- 10. Audacity; Available from: https://www.audacityteam.org/.
- Searfoss AM, Pino JC, Creanza N. Chipper: Open-source software for semiautomated segmentation and analysis of birdsong and other natural sounds. Methods in Ecology and Evolution. 2020;11(4):524–531.

- Markowitz JE, Ivie E, Kligler L, Gardner TJ. Long-range Order in Canary Song. PLOS Computational Biology. 2013;9(5):e1003052. doi:10.1371/journal.pcbi.1003052.
- 13. Jin DZ, Kozhevnikov AA. A compact statistical model of the song syntax in Bengalese finch. PLoS Comput Biol. 2011;7(3):e1001108.
- 14. Berwick RC, Okanoya K, Beckers GJ, Bolhuis JJ. Songs to syntax: the linguistics of birdsong. Trends in cognitive sciences. 2011;15(3):113–121.
- 15. Hedley RW. Complexity, predictability and time homogeneity of syntax in the songs of Cassin's vireo (Vireo cassinii). PloS one. 2016;11(4):e0150822.
- 16. Sober SJ, Brainard MS. Adult birdsong is actively maintained by error correction. Nature neuroscience. 2009;12(7):927.
- Sober SJ, Brainard MS. Vocal learning is constrained by the statistics of sensorimotor experience. Proceedings of the National Academy of Sciences. 2012;109(51):21099–21103.
- Wohlgemuth MJ, Sober SJ, Brainard MS. Linked Control of Syllable Sequence and Phonology in Birdsong. Journal of Neuroscience. 2010;30(39):12936–12949. doi:10.1523/JNEUROSCI.2690-10.2010.
- Aronov D, Andalman AS, Fee MS. A specialized forebrain circuit for vocal babbling in the juvenile songbird. Science. 2008;320(5876):630–634.
- 20. Hahnloser RH, Kozhevnikov AA, Fee MS. An ultra-sparse code underliesthe generation of neural sequences in a songbird. Nature. 2002;419(6902):65–70.
- Gardner TJ, Naef F, Nottebohm F. Freedom and rules: the acquisition and reprogramming of a bird's learned song. Science (New York, NY). 2005;308(5724):1046– 1049. doi:10.1126/science.1108214.
- Burkett ZD, Day NF, Peñagarikano O, Geschwind DH, White SA. VoICE: A semi-automated pipeline for standardizing vocal analysis across models. Scientific reports. 2015;5:10237.
- Daou A, Johnson F, Wu W, Bertram R. A computational tool for automated large-scale analysis and measurement of bird-song syntax. Journal of neuroscience methods. 2012;210(2):147–160.
- 24. Anderson SE, Dave AS, Margoliash D. Template-based automatic recognition of birdsong syllables from continuous recordings. The Journal of the Acoustical Society of America. 1996;100(2):1209–1219.
- 25. Yamahachi H, Zai AT, Tachibana RO, Stepien AE, Rodrigues DI, Cavé-Lopez S, et al. Undirected singing rate as a non-invasive tool for welfare monitoring in isolated male zebra finches. PLOS ONE. 2020;15(8):e0236333. doi:10.1371/journal.pone.0236333.
- Pearre B, Perkins LN, Markowitz JE, Gardner TJ. A fast and accurate zebra finch syllable detector. PLOS ONE. 2017;12(7):e0181992. doi:10.1371/journal.pone.0181992.
- Kogan JA, Margoliash D. Automated recognition of bird song elements from continuous recordings using dynamic time warping and hidden Markov models: A comparative study. The Journal of the Acoustical Society of America. 1998;103(4):2185–2196.

- 28. Troyer lab. Song Browser; Available from: https://www.utsa.edu/troyerlab/ software/SongBrowserManual.pdf.
- 29. Tachibana RO, Oosugi N, Okanoya K. Semi-automatic classification of birdsong elements using a linear support vector machine. PloS one. 2014;9(3):e92584.
- Nicholson D. Comparison of machine learning methods applied to birdsong element classification. In: Proceedings of the 15th Python in Science Conference; 2016. p. 57–61.
- 31. Goodfellow I, Bengio Y, Courville A. Deep learning. MIT press; 2016.
- Farabet C, Couprie C, Najman L, LeCun Y. Learning Hierarchical Features for Scene Labeling. IEEE Transactions on Pattern Analysis and Machine Intelligence. 2013;35(8):1915–1929. doi:10.1109/TPAMI.2012.231.
- 33. Krizhevsky A, Sutskever I, Hinton GE. ImageNet Classification with Deep Convolutional Neural Networks. In: Pereira F, Burges CJC, Bottou L, Weinberger KQ, editors. Advances in Neural Information Processing Systems 25. Curran Associates, Inc.; 2012. p. 1097-1105. Available from: http://papers.nips.cc/paper/ 4824-imagenet-classification-with-deep-convolutional-neural-networks. pdf.
- 34. Graves A. Supervised sequence labelling. In: Supervised sequence labelling with recurrent neural networks. Springer; 2012. p. 5–13.
- Böck S, Schedl M. Polyphonic piano note transcription with recurrent neural networks. In: 2012 IEEE International Conference on Acoustics, Speech and Signal Processing (ICASSP); 2012. p. 121–124.
- Parascandolo G, Huttunen H, Virtanen T. Recurrent Neural Networks for Polyphonic Sound Event Detection in Real Life Recordings. 2016 IEEE International Conference on Acoustics, Speech and Signal Processing (ICASSP). 2016; p. 6440– 6444. doi:10.1109/ICASSP.2016.7472917.
- 37. Graves A, Fernández S, Gomez F, Schmidhuber J. Connectionist temporal classification: labelling unsegmented sequence data with recurrent neural networks. In: Proceedings of the 23rd international conference on Machine learning; 2006. p. 369–376.
- Graves A, Schmidhuber J. Framewise phoneme classification with bidirectional LSTM and other neural network architectures. Neural networks. 2005;18(5-6):602– 610.
- Koumura T, Okanoya K. Automatic recognition of element classes and boundaries in the birdsong with variable sequences. PLoS ONE. 2016;11(7). doi:10.1371/journal.pone.0159188.
- 40. Koumura T. BirdsongRecognition. 2016;doi:10.6084/m9.figshare.3470165.v1.
- Nicholson D, Queen JE, Sober SJ. Bengalese Finch song repository. 2017;doi:10.6084/m9.figshare.4805749.v5.
- Alonso LM, Alliende JA, Goller F, Mindlin GB. Low-dimensional dynamical model for the diversity of pressure patterns used in canary song. Physical Review E. 2009;79(4):041929. doi:10.1103/PhysRevE.79.041929.

- Appeltants D, Gentner TQ, Hulse SH, Balthazart J, Ball GF. The effect of auditory distractors on song discrimination in male canaries (Serinus canaria). Behavioural Processes. 2005;69(3):331–341. doi:10.1016/j.beproc.2005.01.010.
- 44. Alliende J, Lehongre K, Del Negro C. A species-specific view of song representation in a sensorimotor nucleus. Journal of Physiology-Paris. 2013;107(3):193–202. doi:10.1016/j.jphysparis.2012.08.004.
- Ron D, Singer Y, Tishby N. The power of amnesia: Learning probabilistic automata with variable memory length. Machine Learning. 1996;25(2):117–149. doi:10.1007/BF00114008.
- Cohen Y, Shen J, Semu D, Leman DP, Liberti WA, Perkins LN, et al. Hidden neural states underlie canary song syntax. Nature. 2020;582(7813):539–544. doi:10.1038/s41586-020-2397-3.
- 47. Vaswani A, Shazeer N, Parmar N, Uszkoreit J, Jones L, Gomez AN, et al. Attention Is All You Need. arXiv:170603762 [cs]. 2017;.
- Coffey KR, Marx RG, Neumaier JF. DeepSqueak: a deep learning-based system for detection and analysis of ultrasonic vocalizations. Neuropsychopharmacology. 2019;44(5):859–868. doi:10.1038/s41386-018-0303-6.
- Mathis A, Mamidanna P, Cury KM, Abe T, Murthy VN, Mathis MW, et al. DeepLabCut: markerless pose estimation of user-defined body parts with deep learning. Nature Neuroscience. 2018;21(9):1281–1289. doi:10.1038/s41593-018-0209-y.
- Goffinet J, Mooney R, Pearson J. Inferring low-dimensional latent descriptions of animal vocalizations. bioRxiv. 2019; p. 811661. doi:10.1101/811661.
- 51. Sainburg T, Thielk M, Gentner T. Animal Vocalization Generative Network (AVGN): A method for visualizing, understanding, and sampling from animal communicative repertoires. In: CogSci; 2019. p. 3563.
- 52. Sainburg T, Thielk M, Gentner TQ. Latent space visualization, characterization, and generation of diverse vocal communication signals. bioRxiv. 2019; p. 870311.
- 53. Paszke A, Gross S, Chintala S, Chanan G, Yang E, DeVito Z, et al. Automatic differentiation in PyTorch. 2017;.
- 54. Marcel S, Rodriguez Y. Torchvision the machine-vision package of torch. In: Proceedings of the 18th ACM international conference on Multimedia. MM '10. New York, NY, USA: Association for Computing Machinery; 2010. p. 1485–1488. Available from: https://doi.org/10.1145/1873951.1874254.
- Walt Svd, Colbert SC, Varoquaux G. The NumPy Array: A Structure for Efficient Numerical Computation. Computing in Science Engineering. 2011;13(2):22–30. doi:10.1109/MCSE.2011.37.
- Harris CR, Millman KJ, van der Walt SJ, Gommers R, Virtanen P, Cournapeau D, et al. Array programming with NumPy. Nature. 2020;585(7825):357–362.
- 57. Virtanen P, Gommers R, Oliphant TE, Haberland M, Reddy T, Cournapeau D, et al. SciPy 1.0: fundamental algorithms for scientific computing in Python. Nature Methods. 2020;17(3):261–272. doi:10.1038/s41592-019-0686-2.

- 58. Dask Development Team. Dask: Library for dynamic task scheduling; 2016. Available from: https://dask.org.
- 59. team Tpd. pandas-dev/pandas: Pandas; 2020. Available from: https://doi.org/10.5281/zenodo.3509134.
- 60. Hunter JD. Matplotlib: A 2D graphics environment. Computing in Science & Engineering. 2007;9(3):90–95. doi:10.1109/MCSE.2007.55.
- Caswell TA, Droettboom M, Lee A, Hunter J, de Andrade ES, Firing E, et al.. matplotlib/matplotlib: REL: v3.3.2; 2020. Available from: https://doi.org/10. 5281/zenodo.4030140.
- Waskom M, Botvinnik O, Gelbart M, Ostblom J, Hobson P, Lukauskas S, et al.. mwaskom/seaborn: v0.11.0 (Sepetmber 2020); 2020. Available from: https: //doi.org/10.5281/zenodo.4019146.
- Kluyver T, Ragan-Kelley B, Pérez F, Granger BE, Bussonnier M, Frederic J, et al. Jupyter Notebooks-a publishing format for reproducible computational workflows. In: ELPUB; 2016. p. 87–90.
- 64. Hynek Schlawack. attrs;. Available from: https://github.com/python-attrs/ attrs.
- 65. da Costa-Luis C, Larroque SK, Altendorf K, Mary H, Korobov M, Yorav-Raphael N, et al.. tqdm: A fast, Extensible Progress Bar for Python and CLI; 2020. Available from: https://doi.org/10.5281/zenodo.4054194.
- Otchy TM, Wolff SBE, Rhee JY, Pehlevan C, Kawai R, Kempf A, et al. Acute off-target effects of neural circuit manipulations. Nature. 2015;528(7582):358–363. doi:10.1038/nature16442.
- James G, Witten D, Hastie T, Tibshirani R. An introduction to statistical learning. vol. 112. Springer; 2013.