Fast and accurate annotation of acoustic signals with deep neural networks

Elsa Steinfath\textsuperscript{1,2}, Adrian Palacios\textsuperscript{1,2}, Julian R Rottschäfer\textsuperscript{1,2}, Deniz Yuezak\textsuperscript{1,2}, Jan Clemens\textsuperscript{1,3*}

\textsuperscript{1}European Neuroscience Institute - A Joint Initiative of the University Medical Center Göttingen and the Max-Planck-Society, Göttingen, Germany; \textsuperscript{2}International Max Planck Research School and Göttingen Graduate School for Neurosciences, Biophysics, and Molecular Biosciences (GGNB) at the University of Göttingen, Göttingen, Germany; \textsuperscript{3}Bernstein Center for Computational Neuroscience, Göttingen, Germany

\textbf{Abstract} Acoustic signals serve communication within and across species throughout the animal kingdom. Studying the genetics, evolution, and neurobiology of acoustic communication requires annotating acoustic signals: segmenting and identifying individual acoustic elements like syllables or sound pulses. To be useful, annotations need to be accurate, robust to noise, and fast.

We here introduce DAS, a method that annotates acoustic signals across species based on a deep-learning derived hierarchical presentation of sound. We demonstrate the accuracy, robustness, and speed of DAS using acoustic signals with diverse characteristics from insects, birds, and mammals. DAS comes with a graphical user interface for annotating song, training the network, and for generating and proofreading annotations. The method can be trained to annotate signals from new species with little manual annotation and can be combined with unsupervised methods to discover novel signal types. DAS annotates song with high throughput and low latency, allowing realtime annotations for closed-loop experimental interventions.

Overall, DAS is a universal, versatile, and accessible tool for annotating acoustic communication signals.

\textbf{Introduction} Animals produce sounds to foster group cohesion (Haack et al., 1983; Janik and Slater, 1998; Chaverri et al., 2012), to signal the presence of food, friend, or foe (Cäsar et al., 2013; Clay et al., 2012), and to find and evaluate mating partners (Baker et al., 2019; Behr and von Helversen, 2004; Holy and Guo, 2005; Sangiamo et al., 2020). Studying acoustic communication not only provides insight into social interactions within and across species; it can also reveal the mechanisms driving complex behaviors: The genetics and evolution of signal production and recognition (Ding et al., 2016, 2019), the genes and circuits driving song learning (Kollmorgen et al., 2020), or the fast and precise sensorimotor transformations involved in vocal interactions (Coen et al., 2016; Cator et al., 2009; Fortune et al., 2011; Okobi et al., 2019). The first step in many studies of acoustic communication is song annotation: the segmentation and labelling of individual elements in a recording. Acoustic signals are diverse and range from the repetitive long-distance calling songs of crickets, grasshoppers, and anurans (Gerhardt and Huber, 2002), the dynamic and context-specific
courtship songs of vinegar flies or rodents (Coen et al., 2014; Clemens et al., 2018; Neunuebel et al., 2015; Sangiamo et al., 2020), to the complex vocalizations produced by some birds and primates (Lipkind et al., 2013; Weiss et al., 2014; Landman et al., 2020).

This diversity in signal structure has spawned a zoo of annotation tools (Arthur et al., 2013; Coffey et al., 2019; Tachibana et al., 2020; Goffinet et al., 2021; Koumura and Okanoya, 2016; Cohen et al., 2020), but existing methods still face challenges: First, assessing vocal repertoires and their relation to behavioral and neural dynamics (Clemens et al., 2018; Coffey et al., 2019; Neunuebel et al., 2015; Fortune et al., 2011; Okobi et al., 2019) requires annotations to be complete and temporally precise even at low signal levels, but annotation can fail when signals are weak (Coffey et al., 2019; Stern et al., 2017). Second, analyses of large datasets and experimental interventions during behavior (Fortune et al., 2011; Okobi et al., 2019; Bath et al., 2014; Tschida and Mooney, 2012; Stowers et al., 2017) need annotations to be fast, but existing methods are often slow. Last, annotation methods should be flexible and adaptable (Ding et al., 2016, 2019; Clemens et al., 2018; Clemens and Hennig, 2013), but existing methods often work only for restricted types of signals or adapting them to new signals requires tedious manual tuning (Clemens et al., 2018).

In brief, an accurate, fast, and flexible framework for annotating song across species is missing. A general framework would not only improve upon existing methods but would also facilitate the study of species for which automated methods do not yet exist. Deep neural networks have emerged as powerful and flexible tools for solving data annotation tasks relevant for neuroscience such as object recognition, pose tracking, or speech recognition (Krizhevsky et al., 2012; Graves and Jaitly, 2014; Mathis et al., 2018; Pereira et al., 2018; Graving et al., 2019). These methods are not only fast and accurate but also easily adapted to novel signals by non-experts since they only require annotated examples for learning. Recently, deep neural networks have also been used for annotating animal vocalizations (Oikarinen et al., 2019; Coffey et al., 2019; Cohen et al., 2020; Sainburg et al., 2020; Arthur et al., 2021; Goffinet et al., 2021).

We here present a new deep-learning based framework for annotating acoustic signals, called Deep Audio Segmenter (DAS). We test the framework on a diverse set of recordings from insects, birds, and mammals, and show that DAS annotates song in single- and multi-channel recordings with high accuracy. The framework produces annotations with low latency on standard PCs and is therefore ideally suited for closed-loop applications. Small to moderate amounts of manual annotations suffice for adapting the method to a new species and annotation work can be simplified by combining DAS with unsupervised methods. We provide DAS as an open source software package with a graphical user interface for manually annotating audio, training the network, and inferring and proofreading annotations. Integration into existing frameworks for signal analysis or experimental control is possible using a programmatic interface. The code and documentation for DAS are available at https://janclemenslab.org/das/.

Results

Architecture and working principle of DAS

Acoustic signals are defined by features on multiple timescales—the fast harmonic oscillations of the sound carrier (<10 ms), modulations of amplitude (AM) and frequency (FM) (10–100 ms), and the sequencing of different AM and FM patterns into bouts, syllables, or phrases (10–1000 ms). These patterns are typically made explicit using a hand-tuned pre-processing step based on time-resolved Fourier or wavelet transforms (Arthur et al., 2013; Van Segbroeck et al., 2017; Coffey et al., 2019; Oikarinen et al., 2019; Cohen et al., 2020). Most deep-learning based methods then treat this pre-defined spectrogram as an image and use methods derived from computer vision to extract the AM and FM features relevant for annotation (Oikarinen et al., 2019; Coffey et al., 2019; Cohen et al., 2020). Recurrent units are sometimes used to track the sound features over time (Cohen et al., 2020). This approach can produce accurate annotations but has drawbacks: First, the spectrogram constitutes a strong and proven pre-processing step, but it is unsuitable for some
signal types, like short pulsatile signals. Second, the pre-processing step is typically tuned by
hand and may therefore require expert knowledge for it to produce optimal results. Lastly, the
recurrent units used in some methods (Cohen et al., 2020) excel at combining information over
time to provide the context information necessary to annotate spectrally complex signals, but they
can be hard to train and slow to run (Bai et al., 2018).

DAS solves these limitations in three ways: First, the pre-processing step is optional. This makes
DAS more flexible, since signals for which a time-resolved Fourier transform is not appropriate—for
instance, short pulsatile signals—can now also be processed. Second, the optional preprocessing
step is integrated and optimized with the rest of the network. This removes the need to hand-
tune this step and allows the network to learn a preprocessing that deviates from a time-resolved
Fourier or wavelet transform if beneficial (Choi et al., 2017). Integrating the preprocessing into
the network also increases inference speed due to the efficient implementation and hardware
acceleration of deep-learning frameworks. Third and last, DAS learns a task-specific representation
of sound features using temporal convolutional networks (TCNs) (Bai et al., 2018; van den Oord et al.,
2016; Guirguis et al., 2021) (Figure 1–Figure Supplement 1A-E). At the core of TCNs are so-called
dilated convolutions (Yu and Koltun, 2016). In standard convolutions, short templates slide over
the signal and return the similarity with the signal at every time point. In dilated convolutions,
these templates have gaps, allowing to analyze features on longer timescales without requiring
more parameters to specify the template. Stacking dilated convolutions with growing gap sizes
results in a hierarchical, multi-scale representation of sound features, which is ideally suited for
the hierarchical and harmonic structure of animal vocalizations.

The output of the deep neural network in DAS is a set of confidence scores for each audio
sample, corresponding to the probability of each song type (Figure 1C). Annotation labels for the
different song types are mutually exclusive and are produced by comparing the confidence score
to a threshold or by choosing the most probable song type. Brief gaps in the annotations are closed
and short spurious detections are removed to smoothen the annotation. For song types that are
described as events, like the pulses in fly song (Figure 1A), the event times are extracted as local
maxima that exceed a confidence threshold.

DAS accurately annotates song from a diverse range of species

Fly courtship song

We first tested DAS on the courtship song of Drosophila melanogaster, which consists of two major
modes (Figure 1A): The sine song, which corresponds to sustained oscillations with a species-
specific carrier frequency (150 Hz), and two types of pulse song, which consists of trains of short
(5–10 ms) pulses with carrier frequencies between 180 and 500 Hz, produced with a species-specific
interval (35-45 ms in D. melanogaster). Males dynamically choose the song modes based on sen-
sory feedback from the female (Coen et al., 2014, 2016; Clemens et al., 2018; Calhoun et al., 2019).
Despite the relative simplicity of the individual song elements, an accurate annotation of fly song
is challenging because of low signal-to-noise ratio (SNR): The song attenuates rapidly with distance
(Bennet-Clark, 1998) and is highly directional (Morley et al., 2018), which can lead to weak signals
if the male is far from the microphone (Figure 1A). Moreover, the interactions between the flies
introduce pulsatile noise and complicate the accurate and complete annotation of the pulse song.

We first trained DAS to detect the pulse and the sine song recorded using a single microphone
(data from Stern (2014)) and compared the performance of DAS to that of the current state-of-the-
art in fly song segmentation, FlySongSegmenter (FSS) (Arthur et al., 2013; Coen et al., 2014; Clemens
et al., 2018). Annotation performance was quantified using precision, the fraction of correct de-
tections, and recall, the fraction of true song that is detected (Figure 1E, Figure 1–Figure Supple-
ment 1F, G). We counted detected pulses within 10 ms of a true pulse as correct detections. 10 ms
corresponds to 1/4th of the typical interval between pulses in a train and results are robust to the
choice of this value (Figure 1–Figure Supplement 2A). DAS detects pulses with a high precision of
97% — only 3% of all detected pulses are false detections — and a high recall of 96% — it misses
Figure 1. DAS performance for fly song.

A Fly song (black, top) with manual annotations of sine (blue) and pulse (red) song. The spectrogram (bottom) shows the signal's frequency content over time (see color bar).

B DAS builds a hierarchical presentation of song features relevant for annotation using a deep neural network. The output of the network is a confidence score for each sample and song type.

C Confidence scores (top) for sine (blue) and pulse (red) for the signal in A. The confidence is transformed into annotation labels (bottom) based on a confidence threshold (0.5 for sine, 0.7 for pulse). Ground truth (bottom) from manual annotations shown for comparison.

D Confusion matrix for pulse from the test data set. Color indicates the percentage (see color bar) and text labels indicate the number of pulses for each quadrant. All confusion matrices are normalized such that columns sum to 100%. The concentration of values along the diagonal indicates high annotation performance.

E Precision-recall curve for pulse depicts the performance characteristics of DAS for different confidence thresholds (from 0 to 1, black arrow points in the direction of increasing threshold). Recall decreases and precision increases with the threshold. The closer the curve to the upper and right border, the better. The red circle corresponds to the performance of DAS for a threshold of 0.7. The black circle depicts the performance of FlySongSegmenter (FSS) and grey circles the performance of two human annotators.

F Probability density function of temporal errors for all detected pulses (red shaded area), computed as the distance between each pulse annotated by DAS and the nearest manually annotated pulse. Lines depict the median temporal error for DAS (red line, 0.3 ms) and FSS (grey line, 0.1 ms).

G, H Recall of DAS (red line) and FSS (grey line) as a function of the pulse carrier frequency (G) and signal-to-noise ratio (SNR) (H). Red shaded areas show the distributions of carrier frequencies (G) and SNRs (H) for all pulses. DAS outperforms FSS for all carrier frequencies and SNRs.

I Same as in D but for sine. Color indicates the percentage (see color bar) and text labels indicate seconds of sine for each quadrant.

J Same as in E but for sine. The blue circle depicts the performance for the confidence threshold of 0.5.

K Distribution of temporal errors for all detected sine on- and offsets. Median temporal error is 12 ms for DAS (blue line) and 22 ms for FSS (grey line).

L, M Recall for DAS (blue line) and FSS (grey line) as a function of sine duration (L) and SNR (M). Blue shaded areas show the distributions of durations and SNRs for all sine songs. DAS outperforms FSS for all durations and SNRs.

Figure 1–Figure supplement 1. DAS architecture and evaluation.

Figure 1–Figure supplement 2. Performance and the role of context for annotating fly pulse song.

Figure 1–Figure supplement 3. Performance for multi-channel recordings of fly courtship song.
only 4% of all pulses. This is a substantial improvement in recall over FSS, which has slightly higher precision (99%) but misses 13% of all pulses (87% recall) (Figure 1D, E). In DAS, the balance between precision and recall can be controlled via the confidence threshold, which corresponds to the minimal confidence required for labelling a pulse (Figure 1C): Lowering this threshold from 0.7 to 0.5 yields a recall of 99% for pulse song with a modest reduction in precision to 95%. The performance gain of DAS over FSS for pulse stems from better recall at high frequencies (>400Hz) and low SNR (Figure 1G, H). To assess DAS performance for sine song, we evaluated the sample-wise precision and recall. DAS has similar precision to FSS (92% vs 91%) but higher recall (98% vs. 91%) (Figure 1I, J). Recall is higher in particular for short sine songs (<100 ms) and at low SNR (<1.0) (Figure 1L, M). The performance boost for pulse and sine arises because DAS exploits context information, similar to how humans annotate song: For instance, DAS discriminates soft song pulses from pulsatile noise based on the pulse shape but also because song pulses occur in regular trains while noise pulses do not (Figure 1—Figure Supplement 2C). A comparison of DAS’ performance to that of human annotators reveals that our methods exceeds human-level performance for pulse and sine (Figure 1E, J, Table 3).

Temporally precise annotations are crucial, for instance when mapping sensorimotor transformations based on the timing of behavioral or neuronal responses relative to individual song elements (Coen et al., 2014; Srivastava et al., 2017; Long and Fee, 2008; Benichov and Vallentin, 2020). We therefore quantified the temporal error of the annotations produced by DAS. For pulse song, the temporal error was taken as the distance of each pulse annotated by DAS to the nearest true pulse. The median temporal error for pulse is 0.3 ms which is negligible compared to the average duration of a pulse (5-10 ms) or of a pulse interval (35-45 ms) (Deutsch et al., 2019). For sine song, the median temporal error for on- and offsets was 12 ms, which is almost half of that of FSS (22 ms). Sine song can have low SNR (Figure 1M) and fades in and out, making the precise identification of sine song boundaries difficult even for experienced manual annotators (see Figure 1A, C).

Recording song during naturalistic interactions in large behavioral chambers often requires multiple microphones (Coen et al., 2014; Neunuebel et al., 2015). To demonstrate that DAS can process multi-channel audio, we trained DAS to annotate recordings from a chamber tiled with 9 microphones (Coen et al., 2014) (Figure 1—Figure Supplement 3, Figure 1—Figure Supplement 1B). DAS processes multi-channel audio by using filters that take into account information from all channels simultaneously. As is the case for existing methods (Arthur et al., 2013), we achieved maximal performance by training separate networks for the pulse and for the sine song (Table 1). In multi-channel recordings, DAS annotates pulse song with 98% precision and 94% recall, and sine song with 97% precision and 93% recall, and matches the performance of FSS (FSS pulse precision/recall 99/92, sine 95/93) (Figure 1—Figure Supplement 3D-L). Annotations of multi-channel audio have high temporal precision for pulse (DAS 0.3 ms, FSS 0.1 ms) and sine (DAS 8 ms, FSS 15 ms) (Figure 1—Figure Supplement 3E, J). Overall, DAS performs better or as well as the current state-of-the-art method for annotating single and multi-channel recordings of fly song.

Mouse ultrasonic vocalizations
Mice produce ultrasonic vocalizations (USVs) in diverse social contexts ranging from courtship to aggression (Songiamo et al., 2020; Warren et al., 2020; Neunuebel et al., 2015). We tested DAS using audio from an intruder assay, in which an anesthetized female was put into the home cage and the USVs produced by a resident female or male were recorded (Ivanenko et al., 2020). The female USVs from this assay typically consist of pure tones with weak harmonics and smooth frequency modulations that are often interrupted by frequency steps (Figure 2A, B). The male USVs are similar but also contain complex frequency modulations not produced by the females in this assay (Figure 2C, D). Recording noise from animal movement and interaction as well as the frequency steps often challenge spectral threshold-based annotation methods and tend to produce false positive syllables (Tachibana et al., 2020; Coffey et al., 2019). Moreover, weak signals often lead to
Figure 2. DAS performance for mouse ultrasonic vocalizations.

A Waveform (top) and spectrogram (bottom) of USVs produced by a female mouse in response to an anesthetized female intruder. Shaded areas (top) show manual annotations.

B Confidence scores (top) and DAS and manual annotations (bottom) for the female USVs in A. Brief gaps in confidence are filled to ensure smooth annotations.

C Example of male USVs with sex-specific characteristics produced in the same assay.

D Confidence scores (top) and DAS and manual annotations (bottom) for the male USVs in C from a DAS network trained to detect female USVs.

E Confusion matrix from a female-trained network for a test set of female USVs. Color indicates the percentage (see color bar) and text labels the seconds of song in each quadrant.

F Distribution of temporal errors for syllable on- and offsets in female USVs. The median temporal error is 0.3 ms for DAS (brown line) and 0.4 ms for USVSEG (Tachibana et al., 2020), a method developed to annotate mouse USVs (grey line).

G Recall of the female-trained network (brown line) as a function of SNR. The brown shaded area represents the distribution of SNRs for all samples containing USVs. Recall is high even at low SNR.

H Confusion matrix of the female-trained DAS for a test set of male USVs (see C, D for examples). Color indicates the percentage (see color bar) and text labels the seconds of song in each quadrant.

Figure 2—Figure supplement 1. Performance for marmoset vocalizations.

190 missed syllables or imprecisely delimited syllables. We first trained and tested DAS on recordings of a female mouse interacting with an anesthetized female intruder (Figure 2A). DAS annotates the female USVs with excellent precision (98 %) and recall (99 %) (Figure 2E) and low median temporal error (0.3 ms) (Figure 2F). DAS is robust to noise: Even for weak signals (SNR 1/16) the recall is 90 % (Figure 2G). These performance values are on par with that of methods specialized to annotate USVs (Tachibana et al., 2020; Coffey et al., 2019; Van Segbroeck et al., 2017) (see Table 2). USVs of female and male residents have similar characteristics (Figure 2A, B) and the female-trained DAS network also accurately annotated the male vocalizations (Figure 2H). Notably, even the male syllables with characteristics not seen in females in the paradigm were detected (Figure 2D). Overall, DAS accurately and robustly annotates mouse USVs and generalizes across sexes.

Marmoset vocalizations

We next examined the robustness of annotations produced by DAS to noisy recordings and variable vocalization types, by training a network to annotate vocalization from pairs of marmosets (Landman et al., 2020). The recordings contain lots of background noises like faint calls from nearby animals, overdrive from very loud calls of the recorded animals, and large variability within syllables.
ble types (Figure 2-Figure Supplement 1A-D). Recently, a deep-learning based method was shown to produce good performance (recall 77%, precision 85%, 12.5 ms temporal error) when trained on 16000 syllables to recognize seven vocalization types (Oikarinen et al., 2019). We trained DAS on 1/9th of the data (1800 syllables) containing 4 of the 7 vocalization types. Despite the noisy and variable vocalizations, DAS achieves high syllable-wise precision and recall (96%, 92%, (Figure 2-Figure Supplement 1E, F)). Note that DAS obtains this higher performance at millisecond resolution (temporal error 4.4 ms, Figure 2-Figure Supplement 1G), while the method by Oikarinen et al. (2019) only produces annotations with a resolution of 50 ms (Table 2).

Bird song

Bird song is highly diverse and can consist of large, individual-specific repertoires. The spectral complexity and large diversity of the song complicates the annotation of syllable types. Traditionally, syllable types are annotated based on statistics derived from the segmented syllable spectrogram. Recently, good annotation performance has been achieved with unsupervised methods (Sainburg et al., 2020; Goffinet et al., 2021) and deep neural networks (Koumura and Okanoya, 2016; Cohen et al., 2020). We first trained DAS to annotate the song from four male Bengalese finches (data and annotations from Nicholson et al. (2017)). The network was then tested on a random subset of the recordings from all four individuals which contained 37 of the 48 syllable types from the training set (Figure 3A, B, Figure 3-Figure Supplement 1A, B). DAS annotates the bird song with high accuracy: Sample-wise precision and recall are 97% and syllable on- and off-sets are detected with sub-millisecond precision (median temporal error 0.3 ms, Figure 3C). The types of 98.5% the syllables are correctly annotated, with only 0.3% false positives (noise annotated as a syllable), 0.2% false negatives (syllables annotated as noise), and 1% type confusions (Figure 3D, Figure 3-Figure Supplement 1C-D). This results in a low sequence error (corresponding to the minimal number of substitutions, deletions, or insertions required to transform the true sequence of syllables into the inferred one) of 0.012. Overall, DAS performs as well as specialized deep learning-based methods for annotating bird song (Koumura and Okanoya, 2016; Cohen et al., 2020) (Table 2).

To further demonstrate the robustness of DAS, we trained a network to annotate song from Zebra finches. In Zebra finch males, individual renditions of a given syllable type tend to be more variable (Fitch et al., 2002). Moreover, the particular recordings used here (Goffinet et al., 2021) contain background noise from the bird's movement. Despite the variability and noise, DAS annotates the six syllables from a male's main motif with excellent precision and recall, and low temporal error, demonstrating that DAS is robust to song variability and recording noise (Figure 3-Figure Supplement 2).

In summary, DAS accurately and robustly annotates a wide range of signals—from the pulsatile song pulses of flies to the spectrally complex syllables of mammals and birds. DAS therefore constitutes a universal method for annotating acoustic signals that is as good as or better than methods specialized for particular types of signals (Table 2).

**DAS is fast**

To efficiently process large corpora of recordings and to be suitable for closed-loop applications, DAS needs to infer annotations quickly. We therefore assessed the throughput and latency of DAS. Throughput measures the rate at which DAS annotates song and high throughput means that large datasets are processed quickly. Across the five species tested here, DAS has a throughput of 8-82x realtime on a CPU and of 24-267x on a GPU (Figure 4A). This means that a 60-minute recording is annotated in less than 5 minutes on a standard desktop PC and in less than 1.5 minutes using a GPU, making the annotation of large datasets feasible (Figure 4-Figure Supplement 1A-H). The differences in throughput arise from the different sample rates and the network architectures: The marmoset network is fastest because of a relatively shallow architecture with only 2 TCN blocks and a low sampling rate (44.1 kHz). By contrast, the multi-channel *Drosophila* networks have the
Figure 3. DAS performance for the song of Bengalese finches.
A Waveform (top) and spectrogram (bottom) of the song of a male Bengalese finch. Shaded areas (top) show manual annotations colored by syllable type.
B DAS and manual annotation labels for the different syllable types in the recording in A (see color bar). DAS accurately annotates the syllable boundaries and types.
C Confusion matrix for the different syllables in the test set. Color was log-scaled to make rare the annotation errors more apparent (see color bar). Rows depict the probability with which DAS annotated each syllable as any of the 37 types in the test dataset. The type of 98.5% of the syllables were correctly annotated, resulting in the concentration of probability mass along the main diagonal.
D Distribution of temporal errors for the on- and offsets of all detected syllables (green shaded area). The median temporal error is 0.3 ms for DAS (green line) and 1.1 ms for TweetyNet (Cohen et al., 2020), a method developed to annotate bird song (grey line).

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Figure 4. DAS annotates song with high throughput and low latency and requires little data. 

A, B Throughput (A) and latency (B) of DAS (see also Figure 4–Figure Supplement 1). Throughput (A) was quantified as the amount of audio data in seconds annotated in one second of computation time. Horizontal lines in A indicate throughputs of 1, 10, and 40. Throughput is >8x realtime on a CPU (dark shades) and >24x or more on a GPU (light shades). Latency (B) corresponds to the time it takes to annotate a single chunk of audio and is similar on a CPU (dark shades) and a GPU (light shades). Multi-channel audio from flies was processed using separate networks for pulse and sine. For estimating latency of fly song annotations, we used networks with 25 ms chunks, not the 410 ms chunks used in the original network (see Figure 1–Figure Supplement 2).

C Flow diagram of the iterative protocol for fast training DAS.

D Number of manual annotations required to reach 90% of the performance of DAS trained on the full data set shown in Figure 1, Figure 1–Figure Supplement 3, Figure 2, Figure 2–Figure Supplement 1, Figure 3, and Figure 3–Figure Supplement 2 (see also Figure 4–Figure Supplement 3). Performance was calculated as the F1 score, the geometric mean of precision and recall. For most tasks, DAS requires small to modest amounts of manual annotations.

E Current best validation loss during training for fly pulse song recorded on a single channel for 10 different training runs (red lines, 18 minutes of training data). The network robustly converges to solutions with low loss after fewer than 15 minutes of training (40 epochs).

Figure 4–Figure supplement 1. Throughput and latency of inference.

Figure 4–Figure supplement 2. Reducing the chunk duration reduces latency and comes with minimal performance penalties.

Figure 4–Figure supplement 3. DAS requires small to moderate amounts of data for training.

Figure 4–Figure supplement 4. Example of fast training mouse USVs.

Figure 4–Figure supplement 5. DAS performance is robust to changes in the structural parameters of the network.
**DAS requires little manual annotation**

To be practical, DAS should achieve high performance with little manual annotation effort. We find that DAS can be efficiently trained using an iterative protocol (Pereira et al., 2018) (Figure 4C, Figure 4–Figure Supplement 4): Annotate a small set of recordings and train the network for a few epochs; then generate annotations on a larger set of recordings and correct these annotations. Repeat the predict-correct-train cycle on ever larger datasets until performance is satisfactory. To estimate the amount of manual annotations required to achieve high performance, we evaluated DAS trained on subsets of the full training data sets used above (Figure 4–Figure Supplement 3). We then took the number of manual annotations needed to reach 90% of the performance of DAS trained on the full data sets as an upper bound on the data requirements (Figure 4D). With a performance threshold of 90%, the resulting networks will produce sufficiently accurate annotations for creating a larger body of training data with few corrections. Performance was taken as the F1 score, the geometric mean of precision and recall. For single-channel recordings of fly song, fewer than 50 pulses and 20 sine songs are needed to reach 90% of the performance achieved with the full data set. For mouse vocalizations, DAS achieves 90% of its peak performance with fewer than 25 manually annotated syllables. Even for the six syllables from a zebra finch, DAS reaches the 90% threshold with only 48 manually annotated syllables (8 per type). Manually annotating such small amounts of song for flies, mice, or zebra finches takes less than 5 minutes. Likewise, for the song of Bengalese finches, 1-51 (median 8, mean 17) manual annotations are required per syllable type, with one outlier requiring 200 syllables (Figure 4–Figure Supplement 3C). Closer inspection reveals that the outlier results from an annotation error and consists of a mixture of three distinct syllable types (Figure 4–Figure Supplement 3D-F). Even with this outlier, only 626 manually annotated syllables (424 without) are required in total to reach 90% of the test performance of a network trained on >3000 annotated syllables. Data requirements are higher for the multi-channel recordings of fly song (270 pulses and sine songs), and for the noisy and variable marmoset data (1610 annotations, 400 per type), but even in these cases, the iterative training protocol can reduce the manual annotation work.

Overall, DAS requires small to moderate amounts of data for reaching high performance. High throughput (Figure 4A) and small training data sets (Figure 4D) translate to short training times (Figure 4E). The single-channel data sets typically achieve 90% of the performance after less than 10 minutes of training on a GPU. Training on the full data sets typically finishes in fewer than 5 hours. Thus, DAS can be adapted to novel species in short time and with little manual annotation work.

**DAS can be combined with unsupervised methods**

DAS is a supervised annotation method: It discriminates syllable types that have been manually assigned different labels during training. By contrast, unsupervised methods can determine in unlabelled data whether syllables fall into distinct types and if so, classify the syllables (Tabler et al., 2017; Coffey et al., 2019; Clemens et al., 2018; Goffinet et al., 2021; Sainburg et al., 2020; Sangiamo et al., 2020; Arthur et al., 2021). While DAS does not require large amounts of manual annotations (Figure 4D), manual labeling of syllable types can be tedious when differences between syllable types are subtle (Clemens et al., 2018) or when repertoires are large (Sangiamo et al., 2020). In these cases, combining DAS with unsupervised methods facilitates annotation work. To demonstrate the power of this approach, we use common procedures for unsupervised classification, which consist of an initial preprocessing (e.g. into spectrograms) and normalization (e.g. of amplitude) of the syllables, followed by dimensionality reduction and clustering (see Methods) (Clemens et al., 2018; Sainburg et al., 2020; Sangiamo et al., 2020).

For fly song, DAS was trained to discriminate two major song modes, pulse and sine. However, Drosophila melanogaster males produce two distinct pulse types, termed $P_{\text{low}}$ and $P_{\text{fast}}$ (Clemens et al., 2018), and unsupervised classification robustly discriminates the two pulse types as well as the sine song in the DAS annotations (Figure 5A-C). Mouse USVs do not fall into distinct types
We here present Deep Audio Segmenter (DAS), a method for annotating acoustic signals. DAS annotates song in single- and multi-channel recordings from flies (Figure 1, Figure 1–Figure Supplement 3), mammals (Figs Figure 2, Figure 2–Figure Supplement 1), and birds (Figs Figure 3, Figure 3–Figure Supplement 2) accurately, robustly, and quickly (Figure 4A, B). DAS performs as well or better than existing methods that were designed for specific types of vocalizations (Koumura and Okano, 2016; Cohen et al., 2020; Tachibana et al., 2020; Coffey et al., 2019) (Table 2). DAS performs excellently for signals recorded on single and multiple channels (Figure 1, Figure 1–Figure Supplement 3), with different noise levels, and with diverse characteristics. This suggests that DAS is a general method for accurately annotating signals from a wide range of recording setups and species.

Using a user-friendly graphical interface, our method can be optimized for new species without requiring expert knowledge and with little manual annotation work (Figure 4C–E). Network performance is robust to changes in the structural parameters of the network, like filter number and duration, or the network depth (Figure 4–Figure Supplement 5). Thus, the structural parameters do not need to be finely tuned to obtain a performant network for a new species. We have trained networks using a wide range of signal types (Table 4) and these networks constitute good starting points for adapting DAS to novel species. We provide additional advice for the design of novel networks in Methods. This makes the automatic annotation and analysis of large corpora of recordings from diverse species widely accessible.

We show that the annotation burden can be further reduced using unsupervised classification of syllable types, in particular for species with large or individual-specific repertoires (Figure 5) (Clemens et al., 2018; Tabler et al., 2017; Coffey et al., 2019; Goffinet et al., 2021; Sainburg et al., 2020; Arthur et al., 2021). In the future, incorporating recent advances in the self-supervised or semi-supervised training of neural networks will likely further reduce data requirements (Mathis et al., 2021; Raghu et al., 2019; Devlin et al., 2019; Chen and He, 2020). These approaches use unlabelled data to produce networks with a general and rich representation of sound features that can then be fine-tuned for particular species or individuals using few annotated samples. DAS currently does not work well with recordings in which the signals produced by multiple animals overlap. In the future, DAS will be extended with methods for multi-speaker speech recognition to robustly annotate vocalizations from animal groups.

Lastly, the high inference speed (Figure 4A, B) allows integration of DAS in closed-loop systems in which song is detected and stimulus playback or optogenetic manipulation is triggered with low
Figure 5. DAS can be combined with unsupervised methods for song classification.

A Low-dimensional representation obtained using the UMAP (McInnes and Healy, 2018) method of all pulse and sine song waveforms annotated by DAS in a test data set. Data points correspond to individual waveforms and were clustered into three distinct types (colors) using the density-based method HDBSCAN (McInnes et al., 2017).

B, C All waveforms (B) and cluster centroids (C) from A colored by the cluster assignment. Waveforms cluster into one sine (blue) and two pulse types with symmetrical (red, $P_{slow}$) and asymmetrical (orange, $P_{fast}$) shapes.

D, E Low-dimensional representation of the spectrograms of mouse USVs (D) and mean spectrogram for each cluster in D (E). Individual syllables (points) form a continuous space without distinct clusters. Song parameters vary continuously within this space, and syllables can be grouped by the similarity of their spectral contours using k-means clustering.

F, G Low-dimensional representation of the spectrograms of the calls from marmosets (F) and mean spectrogram for each cluster in F (G). Calls separate into distinct types and density-based clustering (colors) produces a classification of syllables that recovers the manual annotations (Figure 4-Figure Supplement 3G, homogeneity score 0.88, completeness score 0.57, v-score 0.69). Most types split into 2 to 4 clusters, reflecting the variability of the call types in marmosets. Colored bars on top of each spectrogram in G correspond to the colors for the individual clusters in F. The dashed line shows the boundary separating trills and twitters.

H, I Low-dimensional representation of the spectrograms of the syllables from one Zebra finch male, mean spectrogram for each cluster in H (I, top), and example of each clustered syllable within the motif (I, bottom). Density-based clustering (colors) recovers the 6 syllable types forming the male’s motif. Colored bars on top of each spectrogram in I correspond to the colors for the individual clusters in H.

J, K Low-dimensional representation of the spectrograms of the syllables from 4 Bengalese finch males (J) and mean spectrogram for each cluster in J (K). Syllables separate into distinct types and density-based clustering (colors) produces a classification of syllables that closely matches the manual annotations (homogeneity score 0.96, completeness score 0.89, v-score 0.92).

X-axes of the average spectrograms for each cluster do not correspond to linear time, since the spectrograms of individual syllables were temporally log-rescaled and padded prior to clustering. This was done to reduce the impact of differences in duration between syllables.
latency (Bath et al., 2014; Stowers et al., 2017). In combination with realtime pose tracking (Mathis et al., 2018; Pereira et al., 2018; Graving et al., 2019), DAS provides unique opportunities to tailor optogenetic manipulations to specific behavioral contexts, for instance to dissect the neural circuits underlying acoustic communication in interacting animals (Coen et al., 2014; Fortune et al., 2011; Okobi et al., 2019).

**Methods and Materials**

Instructions for installing and using DAS can be found at https://janclemenslab.org/das. Code for the das python module is available at https://github.com/janclemenslab/das, code for the unsupervised methods is at https://github.com/janclemenslab/das_unsupervised. All fitted models (with example data and code) can be found at https://github.com/janclemenslab/das-menagerie.

We also provide instructions for training DAS using google colab, which provides a GPU-accelerated python environment. Colab removes the need to install GPU libraries: Annotations can be made locally in the GUI without a GPU and training and prediction are done on GPU-accelerated nodes in the cloud. See this notebook for details: http://janclemenslab.org/das/tutorials/colab.html.

**Data sources**

All data used for testing DAS were published previously. Sources for the original data sets, for the data and annotations used for training and testing, and for the fitted models are listed in Table 5. Single-channel recordings of Drosophila melanogaster (strain OregonR) males courting females were taken from Stern (2014). The multi-channel data from Drosophila melanogaster (strain NM91) males courting females were recorded in a chamber tiled with nine microphones (Coen et al., 2014) and was previously published in Clemens et al. (2018). Annotations for fly song were seeded with FlySongSegmenter (Arthur et al., 2013; Coen et al., 2014) and then manually corrected. Recordings of mouse USVs were previously published in Ivanenko et al. (2020). The USVs were manually labelled using the DAS graphical interface. Marmoset recordings were taken from the data published with Landman et al. (2020). Since we required more precise delineation of the syllable boundaries than was provided in the published annotations, we manually fixed annotations for a subset of the data that then was used for training and testing. The network was trained and tested on a subset of 4 vocalization types (eks/trills/tsiks/twitters, N=603/828/115/868). The remaining vocalization types were excluded since they had 60 or fewer instances in our subset. To test DAS on bird songs, we used a publicly available, hand-labeled collection of song from four male Bengalese finches (Nicholson et al., 2017) and recordings of female-directed song from a male Zebra finch from Goffinet et al. (2021). For training and testing the Zebra finch network, we manually labelled 473 syllables of 6 types (320 seconds of recordings).

**DAS network**

DAS is implemented in Keras (Chollet et al., 2015) and Tensorflow (Abadi et al., 2016). At its core, DAS consists of a stack of temporal convolutional blocks, which transform an input sequence of audio data into an output sequence of labels.

**Inputs**

DAS takes as input raw, single or multi-channel audio. Pre-processing of the audio using a Wavelet or short-time Fourier transform is optional and integrated into the network. DAS processes audio in overlapping chunks (Figure 1–Figure Supplement 1A-D). The chunking accelerates annotations since multiple samples are annotated in a single computational step. Edge effects are avoided by processing overlapping chunks and by discarding a number of samples at the chunk boundaries. The overlap depends on the number of layers and the duration of filters in the network.
The trainable short-term Fourier transform (STFT) frontend is an optional step and was implemented using kapre (Choi et al., 2017). Each frequency channel in the output of the frontend is the result of two, one-dimensional strided convolutions which are initialized with the real and the imaginary part of discrete Fourier transform kernels:

\[
x(i, f) = \sum_{\tau=0}^{T-1} x(i \ast s + \tau)[\cos(2\pi f \tau / T) - i \sin(2\pi f \tau / T)]
\]

\[
y(i, f) = \log_{10}(\sqrt{\Re x(i, f)^2 + \Im x(i, f)^2})
\]

where \( f \) is the frequency, \( s \) is the stride, and \( T \) is the filter duration. The stride results in downsampling of the input by a factor \( s \).

The STFT kernels are optimized with all other parameters of the network during training. The STFT frontend was used for mammal and bird signals, but not for fly song. In the mammal and bird networks, we used 33 STFT filter pairs with a duration \( T = 64 \) samples and a stride \( s = 16 \) samples. For mouse and bird song, the STFT frontend sped up training and inference, and had a small positive impact on performance. For fly song, the STFT frontend tended to reduce performance and was omitted.

Temporal convolutional blocks

Temporal convolutional network (TCN) blocks are central to DAS and produce a task-optimized hierarchical representation of sound features at high temporal resolution (Bai et al., 2018). Each TCN block consists of a stack of so-called residual blocks (Figure 1–Figure Supplement 1E) (He et al., 2016):

A dilated convolutional layer filters the input with a number of kernels of a given duration: \( y_i(t) = \sum_{\tau, \gamma} k_i(\tau, \gamma) \ast x(t - a \tau, \gamma) \), where \( k_i(\tau, \gamma) \) is the kernel, \( x(t, \gamma) \) is the input on channel \( \gamma \) at time \( t \), \( y_i(t) \) the output and \( a \) the gap or skip size (Yu and Koltun, 2016). In old-fashioned convolution \( a = 1 \).

Increasing \( a \) allows the kernel to span a larger range of inputs with the same number of parameters and without a loss of output resolution. The number of parameters is further reduced for networks processing multi-channel audio, by using separable dilated convolutions in the first two TCN blocks (Mamalet and Garcia, 2012). In separable convolutions, the full two-dimensional \( k(\tau, \gamma) \) convolution over times and channels is decomposed into two one-dimensional convolutions. First, a temporal convolution, \( k_t(\tau, 1) \), is applied to each channel and then \( N \) channel convolutions, \( k^\gamma(1, \gamma) \), combine information across channels. Instead of \( \tau \times \gamma \) parameters, the separable convolution only requires \( \tau + N \times \gamma \) parameters. Note that each temporal convolution is applied to each channel, leading to a sharing of filter parameters across channels. This makes explicit the intuition that some operations should be applied to all channels equally. We also tested an alternative implementation, in which individual channels were first processed separately by a single-channel TCN, the outputs of the TCN blocks for each channel were concatenated, and then fed into a stack of standard TCNs with full two-dimensional convolutions. While this architecture slightly increased performance it was also much slower and we therefore chose the architecture with separable convolutions. Architecture choice ultimately depends on speed and performance requirements of the annotation task.

A rectifying linear unit transmits only the positive inputs from the dilated convolutional layer by setting all negative inputs to 0: \( y_i = \max(0, y_i) \).

A normalization layer rescales the inputs to have a maximum absolute value close to 1: \( y_i / (\max(|y_i|) + 10^{-5}) \).

The output of the residual block, \( z(t) \), is then routed to two targets: First, it is added to the input: \( o(t) = x(t) + z(t) \) and fed into subsequent layers. Second, via a so-called skip connection, the outputs of all residual blocks are linearly combined to produce the network’s final output (van den Oord et al., 2016).
A single TCN block is composed of a stack of five residual blocks. Within a stack, the skip size doubles — from 1 in the first to $2^5 = 16$ in the final residual block of a stack. This exponential increase in the span of the filter $\tau \ast a$ allows the TCN block to produce a hierarchical representation of its inputs, from relatively low-level features on short timescales in early stacks to more derived features on longer timescales in late stacks. Finally, a full network consists of a stack of 2 to 4 TCN blocks, which extract ever more derived features (Figure 1–Figure Supplement 1A-D).

Output

The network returns a set confidence scores—one for each song type (and for no song)—for each sample from a linear combination of the output of each residual block in the network, by using a single dense layer and a softmax activation function. Re-using information from all blocks via so-called skip connections ensures that downstream layers can discard information from upstream layers and facilitates the generation of specialized higher-order presentations. If the input recording got downsampled by a STFT frontend, a final upsampling layer restores the confidence scores to the original audio rate by repeating values. The parameters of all used networks are listed in Table 4.

Choice of structural network parameters

DAS performance is relatively robust to the choice of structural network parameters like filter duration and number, or network depth (Figure 4–Figure Supplement 5). The networks tested here are good starting points for adapting DAS to your own data (Table 4). In our experience, a network with 32 filters, filter duration 32 samples, 3 TCN blocks, and a chunk duration of 2048 samples will produce good results for most signals. A STFT-downsampling layer with 32 frequency band and 16x downsampling should be included for most signals except when the signals have a pulsatile character. These parameters have been set as defaults when creating a new DAS network. Given that DAS trains quickly (Figure 4E), network structure can be optimized by training DAS networks over a grid of structural parameters, for instance to find the simplest network (in terms of the number of filters and TCN blocks) that saturates performance but has the shortest latency. We here provide additional guidelines for choosing a network’s key structural parameters:

The chunk duration corresponds to the length of audio the network processes in one step and constitutes an upper bound for the context available to the network. Choose chunks sufficiently long so that the network has access to key features of your signal. For instance, for fly song, we ensured that a single chunk encompasses several pulses in a train, so the network can learn to detect song pulses based on their regular occurrence in trains. Longer chunks relative to this timescale can reduce short false positive detections, for instance for fly sine song and for bird song. Given that increasing chunk duration does not increase the number of parameters for training, we recommend using long chunks unless low latency is of essence (see below).

Downsampling/STFT weakly affects performance but strongly accelerates convergence during training. This is because A) the initialization with STFT filters is a good prior that reduces the number of epochs it takes to learn the optimal filters, and B) the downsampling reduces the data bandwidth and thereby the time it takes to finish one training epoch. The overall increase in performance from adding the STFT layer is low because convolutional layers in the rest of the network can easily replicate the computations of the STFT layer. For short pulsatile signals or signals with low sampling rates, STFT and downsampling should be avoided since they can decrease performance due to the loss of temporal resolution.

The number of TCN blocks controls the network’s depth. A deeper network can extract more high-level features, though we found that even for the spectro-temporally complex song of Bengalese finches, deeper networks only weakly improved performance (Figure 4–Figure Supplement 5).

Multi-channel audio can be processed with multi-channel filters via full convolutions or with shared channel-wise filters via time-channel separable convolutions. This can be set on a per-TCN-block basis. We recommend to use separable convolutions in the first 1–2 layers, since basic
feature extraction is typically the same for each channel. Later layers can then have full multi-channel filters to allow more complex combination of information across channels.

**Real-time performance** can be optimized by reducing networks complexity and chunk duration (Figure 4–Figure Supplement 2). We recommend starting with the default parameters suggested above and then benchmarking latency. If required, latency can then be further reduced by reducing chunk duration, the number and duration of filters, and the number of TCN blocks.

**Training**

Networks were trained using the categorical cross-entropy loss and the Adam optimizer (Kingma and Ba, 2015) with a batch size of 32. Prediction targets were generated from fully annotated recordings and one-hot-encoded: Segments were coded as binary vectors, with \( y_i(t) = 1 \) if a segment of type \( i \) occurred at time \( t \), and \( y_i(t) = 0 \) otherwise. To encode uncertainty in the timing of fly song pulses, the pulses were represented as Gaussian bumps with a standard deviation of 1.6 ms. A “no song” type was set to \( y_{\text{no song}}(t) = 1 - \sum y_i(t) \). That way, \( y \) corresponds to the probability of finding any of the annotated song types or no song. For bird song, short gaps (6.25 ms, 200 samples at 32 kHz) were introduced between adjacent syllables to aid the detection of syllable on- and offsets after inference. That way, syllable on- and offsets could be unequivocally detected as changes from “no song” to any of the syllables. This reduced the amount of false positive on- and offsets from switches in the label within a syllable.

Typically, multiple fully annotated recordings were combined in a data set. Each recording was split 80:10:10 into a training, validation, and test set. The validation and test data were randomly taken from the first, the middle or the last 10% of each recording. Given the uneven temporal distribution of call types in the marmoset recordings, we split the data 60:20:20 to ensure that each call type was well represented in each split. For all networks, training was set to stop after 400 epochs or earlier if the validation loss was not reduced for at least 20 epochs. Training typically stopped within 40-80 epochs depending on the dataset. The test set was only used after training, for evaluating the model performance.

**Generation of annotations from the network output**

The confidence scores produced by the model correspond to the sample-wise probability for each song type. To produce an annotation label for each sample, the confidence scores were further processed to extract event times and syllable segments. In the resulting annotations, song types are mutually exclusive, that is, each sample is labelled as containing a single song type even if song types overlap.

Event times for event-like song types like fly pulse song were determined based on local maxima in the confidence score, by setting a threshold value between 0 and 1 and a minimal distance between subsequent peaks (using peakutils (Negri and Vestri, 2017)). For the pulse song of flies, we set a minimal distance of 10 ms and a threshold of 0.7 for single channel data (Figure 1) and 0.5 for multi-channel data (Figure 1–Figure Supplement 3).

For segment-like song types like fly sine song or the syllables of mouse, marmoset, and bird song, we first transformed the sample-wise probability into a sequence of labels using argmax, \( y(i, t) \). The resulting annotation of segments was then smoothed by filling short gaps (flies 20 ms, mice 10 ms, marmosets and birds 5 ms) and removing very short detections (flies 20 ms, mice 5 ms, marmosets and birds 30 ms). These values were chosen based on the statistics of song found in the training data. Syllable on- and offsets were detected as changes from no-song to song and song to no-song, respectively. For bird and marmoset vocalizations, syllable labels were determined based on a majority vote, by calculating the mode of the sample-wise labels for each detected syllable.

**Evaluation**

DAS was evaluated on segments of recordings that were not used during training.
Events

For events—fly pulse song, or the on- and offsets of segments—we matched each true event with its nearest neighbor in the list of true events and counted as true positives only events within a specified distance from a true event. For the pulse song of flies as well as for the onsets and offsets of mouse, marmoset, and bird syllables, this distance was set to 10 ms. Results were robust to the specific choice of the distance threshold (Figure 1–Figure Supplement 2A). For the onsets and offsets of fly sine song and of the marmoset vocalizations, we set this distance to 40 ms, since these signals tended to fade in and out, making the delineation of exact boundaries difficult. False positive events were counted if the distance from a detected event to the nearest true event exceeded the distance threshold or if another detected event was closer to each true event within the distance threshold. If several detected pulses shared the same nearest true pulses, only the nearest of those was taken as a true positive, while the remaining detections were matched with other true pulses within the distance threshold or counted as false positives.

False negatives were counted as all true events without nearby detected events. For pulse, pseudo true negative events were estimated as the number of tolerance distances (2x tolerance distance) fitting into the recording, minus the number of pulses. These true negatives for pulse do not influence precision, recall, and F1-scores and are only used to fill the confusion matrices in Figure 1D, I and Figure 1–Figure Supplement 3C, H. Pulse and sine song were evaluated only up to the time of copulation.

Matching segment labels

For songs with only one syllable type, we compared the predicted and true labels for each sample to compute the confusion matrix (Figure 1–Figure Supplement 1F, G). In the case of multiple syllable types, the mode of the true and predicted labels for the samples of each detected syllable were compared. A true positive was counted if the mode of the true labels was the same for the samples covered by the detected syllable. Using the true syllables as reference produces similar results (Figure 2–Figure Supplement 1E, F and Figure 3–Figure Supplement 1D, E).

Performance scores

From the false negative (FN), false positive (FP), and true positive (TP) counts we extracted several scores: Precision (P)—the fraction of true positive out of all detections TP/(FP+TP)—and recall (R)—the fraction of true positives out of all positives TP/(TP+FN). The F1 score combines precision and recall via their geometric mean: $2 \times P \times R/(P + R)$. For datasets with many different syllable types, we also used as a summary measure of performance the accuracy—the fraction of correctly labelled syllables: $(TP+TN)/(TP+TN+FP+FN)$. For comparison with other studies, we additionally provide the error rate for the song of Bengalese finches, which is based on the Levenshtein edit distance and corresponds to the minimal number of inserts, deletions, and substitutions required to transform the sequence of true syllable labels into the sequence of predicted syllable labels normalized by the length of the true sequence (Koumura and Okanoya, 2016; Cohen et al., 2020).

Temporal precision

The temporal precision for events (pulses, syllable onsets and offsets) was calculated as the median absolute distance between all matched events.

Annotation errors for Bengalese finches

The network for annotating bird song was trained on all syllable types. We removed from the test data one syllable type with only a single instance in the test set (which was correctly classified), because the performance could not be assessed reliably based on a single instance. We also excluded as annotation error a syllable type that contained syllables of more than 6 distinct types.
Estimation of signal-to-noise ratio from audio recordings

To assess the robustness of annotation performance to noise, we assessed the recall of DAS for epochs with different signal-to-noise ratios (SNRs) for the fly and the mouse networks. Because of fundamental differences in the nature of the signals, SNR values were computed with different methods and are therefore not directly comparable across species.

Pulse

Pulse waveforms were 20 ms long and centered on the peak of the pulse energy. The root-mean square (RMS) amplitudes of the waveform margins (first and last 5 ms) and center (7.5-12.5 ms) were taken as noise and signal, respectively. RMS is defined as $\sqrt{\sum x(i)^2}$. For multi-channel recordings, the pulse waveform from the channel with the highest center RMS was chosen to calculate the SNR.

Sine

Signal was given by the RMS amplitudes of the recording during sine song. Noise is the RMS amplitude in the 200 ms before and after each sine song, with a 10 ms buffer. For instance, if a sine song ended at 1000 ms, the recording between 1010 and 1210 ms was taken as noise. From the 200 ms noise, we excluded samples that were labelled as sine or pulse and included intervals between pulses. For multi-channel recordings, the SNR was calculated for the channel with the largest signal amplitude.

Mouse

We assumed an additive noise model: $\sigma_{total}^2 = \sigma_{signal}^2 + \sigma_{noise}^2$. $\sigma^2$ is the squared signal averaged over a window of 1 ms. Since noise variance changed little relative to the signal variance in our recordings, we can assume constant noise over time to calculate the signal strength: $\sigma_{signal}^2 = \sigma_{total}^2 - \sum \sigma_{noise}^2$. The sample-wise SNR is then given by $SNR(i) = \sigma_{signal}(i)^2 / \sum \sigma_{noise}^2$.

Speed benchmarks

Inference speed was assessed using throughput and latency. Throughput is the number of samples annotated per second and latency is the time it takes to annotate a single chunk. Throughput and latency depend on the chunk duration—the duration of a recording snippet processed by the network at once—and on the batch size—the number of chunks processed during one call. Larger batches maximize throughput by more effectively exploiting parallel computation in modern CPUs and GPUs and reducing overheads from data transfer to the GPU. This comes at the cost of higher latency, since results are available only after all chunks in a batch have been processed. Using small batch sizes and short chunks therefore reduces latency, since results are available earlier, but this comes at the cost of reduced throughput because of overhead from data transfer or under-utilized parallel compute resources. To assess throughput and latency, run times of model.predict were assessed for batch sizes ranging from 1 to 1024 (log spaced) with 10 repetitions for each batch size after an initial warmup run (Figure 4-Figure Supplement 1A-L). Results shown in the main text are from a batch size corresponding to 1s of recording for throughput (Figure 4A) and a batch size of 1 for latency (Figure 4B, see also Figure 4-Figure Supplement 1). For fly song, latency was optimized by reducing the chunk size to 25.6 ms (Figure 4-Figure Supplement 2). Benchmarks were run on Windows 10, Tensorflow 2.1, with the network either running on a CPU (Intel i7-7770, 3.6GHz) or on a GPU (GTX1050 Ti 4GB RAM).

We also benchmarked the throughput of existing methods for comparison with DAS (Table 2). Since neither of the methods considered are designed to be used in ways in which latency can be fairly compared to that of DAS, we did not assess latency. The throughput values include all pre-processing steps (like calculation of a spectrogram) and comparisons to DAS were done using the same hardware (CPU for FSS and USVSEG, GPU for TweetyNet and Oikarinen et al. (2019)).
throughput of FSS (Arthur et al., 2013; Coen et al., 2014) was tested using 400 seconds of single-channel and 9-channel recordings in Matlab2019a. USVSEG (Tachibana et al., 2020) was tested on a 72 second recording in Matlab2019a. TweetyNet (Cohen et al., 2020) was tested using a set of 4 recordings (total duration 35 seconds). Throughput for TweetyNet was given by the combined run-times of the pre-processing steps (calculating of spectrograms from raw audio and saving them as temporary files) and the inference steps (running the network on a GPU). For the previously published network for annotating marmoset calls (Oikarinen et al., 2019), we relied on published values for estimating throughput: A processing time of 8 minutes for a 60 minute recording corresponds to a throughput of 7.5 s/s.

Data economy

For estimating the number of manual annotations required to train obtain accurate annotations, we trained the networks using different fractions of the full training and validation sets (for instance, 0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1.0). Performance of all networks trained on the different subsets was evaluated on the full test set. The number of manual annotations in each subset was determined after training from the training and validation sets. The number of annotations required to exceed 90% of the F1 score of a model trained on the full data sets was calculated based on a lowess fit (Cleveland, 1979) to the data points (Figure 4A, B).

Unsupervised classification

Segmented signals were clustered using unsupervised methods described previously in Clemens et al. (2018), Sainburg et al. (2020), and Sangiamo et al. (2020). First, signals were pre-processed: For fly song, pulse and sine waveforms of duration 15 ms were extracted from the recording, aligned to their peak energy, normalized to unit norm, and adjusted for sign (see Clemens et al. (2018) for details). For mouse, marmoset, and bird vocalizations, we adapted the procedures described in Sainburg et al. (2020): Noise was reduced in the bird song recordings using the noisereduce package (https://github.com/timsainb/noisereduce). For mouse and marmoset vocalizations, noise reduction tended to blur the spectral contours and was omitted. Then, syllable spectrograms were extracted from mel spectrograms of the recordings. The noise floor of the spectrogram at each frequency was estimated as the median spectrogram over time and each spectral band was then divided by the frequency-specific noise floor value. Finally, the spectrogram values were log-transformed and thresholded at 0 for mice and 2 for marmosets and birds after visual inspection of the spectrograms to further remove background noise. To reduce differences in the duration of different syllables, all syllables were first log resized in time (scaling factor 8) and then padded with zeros to the duration of the longest syllable in the data set. Lastly, the frequency axis of the spectrograms for mouse syllables were aligned to the peak frequency, to make clustering robust to jitter in the frequency of the thin spectral contours (Sangiamo et al., 2020). The peak frequency of each mouse syllable was calculated from its time-averaged spectrogram, and only the 40 spectrogram frequencies around the peak frequency were retained. The dimensionality of the pre-processed waveforms (fly) or spectrograms (mouse, marmoset, birds) was then reduced to two using the UMAP method (McInnes and Healy, 2018) (mindist=0.5, 0.1 for marmosets to improve separation of clusters). Finally, signals were grouped using unsupervised clustering. For the fly, marmoset, and bird signals, the UMAP distribution revealed distinct groups of syllables and we used a density-based method to cluster the syllables (Campello et al. (2013), min_samples=10, min_cluster_size=20). For mouse USVs, no clusters were visible in the UMAP distribution and density-based clustering failed to identify distinct groups of syllables. Syllables were therefore split into 40 groups using k-means clustering.

Open source software used

- avgn https://github.com/timsainb/avgn_paper (Sainburg et al., 2020)
- hdbscan (McInnes et al., 2017)
• ipython (Pérez and Granger, 2007)
• jupyter (Kluyver et al., 2016)
• kapre (Choi et al., 2017)
• keras (Chollet et al., 2015)
• keras-tcn https://github.com/philipperemy/keras-tcn
• librosa (McFee et al., 2015)
• matplotlib (Hunter, 2007)
• noisereduce https://github.com/timsainb/noisereduce
• numpy (Harris et al., 2020)
• pandas (McKinney, 2010)
• peakutils (Negri and Vestri, 2017)
• scikit-learn (Pedregosa et al., 2011)
• scipy (Virtanen et al., 2020)
• seaborn (Waskom et al., 2017)
• snakemake (Köster and Rahmann, 2018)
• tensorflow (Abadi et al., 2016)
• UMAP (McInnes and Healy, 2018)
• zarr (Miles et al., 2020)
• xarray (Hoyer and Hamman, 2017)

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Table 1. Precision, recall, and temporal error of DAS.
Precision and recall values are sample-wise for all except fly pulse song, for which it is event-wise. The number of classes includes the “no song” class. (p) Pulse, (s) Sine.

| Species              | Trained                  | Classes | Threshold | Precision [%] | Recall [%] | Temporal error [ms] |
|----------------------|--------------------------|---------|-----------|---------------|------------|---------------------|
| Fly single channel   | Pulse (p) & sine (s)     | 3       | 0.7       | 97/92 (p/s)   | 96/98 (p/s)| 0.3/12 (p/s)        |
| Fly multi channel    | Pulse (p)                | 2       | 0.5       | 98            | 94         | 0.3                 |
| Fly multi channel    | Sine (s)                 | 2       | 0.5       | 97            | 93         | 8                   |
| Mouse                | Female                   | 2       | 0.5       | 98            | 99         | 0.3                 |
| Marmoset             | 5 male-female pairs      | 5       | 0.5       | 85            | 91         | 4.4                 |
| Bengalese finch      | 4 males                  | 49 (38 in test set) | 0.5 | 97 | 97 | 0.3 |
| Zebra finch          | 1 males                  | 7       | 0.5       | 98            | 97         | 1.2                 |

Table 2. Comparison to alternative methods.
Methods used for comparisons:
(1) Arthur et al. (2013), (2) Tachibana et al. (2020), (3) Oikarinen et al. (2019), (4) Cohen et al. (2020).
(A,B) DAS was trained on 1825/15970 syllables which contained 4/7 of the call types from Oikarinen et al. (2019).
(B) The method by Oikarinen et al. (2019) produces an annotation every 50 ms of the recording - since the on/offset can occur anywhere within the 50 ms, the expected error of the method by Oikarinen et al. (2019) is at least 12.5 ms.
(C) The method by Oikarinen et al. (2019) annotates 60 mins of recordings in 8 minutes.
(D) Throughput assessed on the CPU, since the methods by Arthur et al. (2013) and Tachibana et al. (2020) do not run on a GPU.
(E) Throughput assessed on the GPU. The methods by Cohen et al. (2020) and Oikarinen et al. (2019) use a GPU.

| Species              | Precision [%] | Recall [%] | Jitter [ms] | Throughput [s/s] |
|----------------------|--------------|------------|-------------|------------------|
| Fly single (1)       | DAS 97/92 (p/s) | other 99/91 | 0.3/12 (p/s) | 15 (4D)         |
| Fly multi (1)        | DAS 98 | 99/96 (p/s) | 87/91 | 0.1/22 | 8 (p) 0.4 (p+s)(D) |
| Fly multi (1)        | DAS 97 | 95/93 | 93/8.0 | 15.0 | 8 (s) 0.4 (p+s)(D) |
| Mouse (2)            | DAS 98 | 98/99 | 99/0.3 | 0.4 | 12 (4D) |
| Marmoset (3)         | DAS 96 | 85(A) | 92/77(A) | 4.4 | 12.5(B) 7.5 (C,E) |
| Bengalese finch (4)  | DAS 99 | 99/99 | 99/0.3 | 1.1 | 15 (5 (E) |
| Zebra finch (4)      | DAS 100 | 100/100 | 100/100 | 1.3 | 2.0 18 (5 (E) |

References
Abadi M, Barham P, Chen J, Chen Z, Davis A, Dean J, Devin M, Ghemawat S, Irving G, Isard M, Kudlur M, Levenberg J, Monga R, Moore S, Murray DG, Steiner B, Tucker P, Vasudevan V, Warden P, Wicke M, et al. TensorFlow: A System for Large-Scale Machine Learning. In: OSDI’16, USA: USENIX Association; 2016. p. 265–283.

Arthur BJ, Ding Y, Sosale M, Khalif F, Kim E, Waddell P, Turaga SC, Stern DL. SongExplorer: A deep learning workflow for discovery and segmentation of animal acoustic communication signals. bioRxiv. 2021; doi: 10.1101/2021.03.26.437280.

Arthur BJ, Sunayama-Morita T, Coen P, Murthy M, Stern DL. Multi-channel acoustic recording and automated analysis of Drosophila courtship songs. BMC Biology. 2013; 11(1):11.

Table 3. Comparison to human annotators for fly song.
See also Figure 1E, J.

| Annotator | Sine recall [%] | Sine precision [%] | Pulse recall [%] | Pulse precision [%] |
|-----------|----------------|-------------------|------------------|-------------------|
| Human A   | 89             | 98                | 99               | 93                |
| Human B   | 93             | 91                | 98               | 88                |
| FSS       | 91             | 91                | 87               | 99                |
| DAS       | 98             | 92                | 96               | 97                |
Table 4. Structural parameters of the tested networks.

| Species                | Rate [kHz] | Chunk [samples] | Channels | STFT downsample | Separable conv. | TCN stacks | Kernel size [samples] | Kernels |
|------------------------|------------|-----------------|----------|-----------------|-----------------|------------|-----------------------|---------|
| Fly single channel     | 10.0       | 4096            | 1        | -               | -               | 3          | 32                    | 32      |
| Fly multi channel (pulse) | 10.0     | 2048            | 9        | -               | TCN stacks 1+2  | 4          | 32                    | 32      |
| Fly multi channel (sine) | 10.0     | 2048            | 9        | -               | TCN stacks 1+2  | 4          | 32                    | 32      |
| Mouse                  | 300.0      | 8192            | 1        | 16x             | -               | 2          | 16                    | 32      |
| Marmoset               | 44.1       | 8192            | 1        | 16x             | -               | 2          | 16                    | 32      |
| Bengales finch         | 32.0       | 1024            | 1        | 16x             | -               | 4          | 32                    | 64      |
| Zebra finch            | 32.0       | 2048            | 1        | 16x             | -               | 4          | 32                    | 64      |

Table 5. Sources of all data used for testing DAS.

"Data" refers to the data used for DAS and to annotations that were either created from scratch or modified from the original annotations (deposited under https://data.goettingen-research-online.de/dataverse/das).

"Original data" refers to the recordings and annotations deposited by the authors of the original publication.

"Model" points to a directory with the model files as well as a small test data set and demo code for running the model (deposited under https://github.com/janclemenslab/das-menagerie).

| Species                | Reference              | Data and model repositories                      |
|------------------------|------------------------|---------------------------------------------------|
| Fly single channel     | *Stern (2014)*         | data: https://doi.org/10.25625/TP4ODR             |
|                        |                        | original data: https://www.janelia.org/lab/stern-lab/tools-reagents-data |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/dmel_single |
| Fly multi channel      | *Clemens et al. (2018)* | data: https://doi.org/10.25625/8KAKHJ         |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/dmel_multi |
| Mouse                  | *Ivanenko et al. (2020)* | data: https://doi.org/10.25625/VVSJKCH |
|                        |                        | original data: https://data.donders.ru.nl/collections/dcn/DSC_620840_0003_891 |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/mouse |
| Marmoset               | *Landman et al. (2020)* | data: https://doi.org/10.25625/DYG3KV         |
|                        |                        | original data: https://osf.io/q4bm3/          |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/marmoset |
| Bengales finch         | *Nicholson et al. (2017)* | data: https://doi.org/10.25625/ENKMGJS       |
|                        |                        | original data: https://doi.org/10.6084/m9.figshare.4805749.v6 |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/bengalese_finch |
| Zebra finch            | *Goffinet et al. (2021)* | data: https://doi.org/10.25625/ZXJJYY        |
|                        |                        | original data: https://research.repository.duke.edu/concern/datasets/9k41zf38g |
|                        |                        | model: https://github.com/janclemenslab/das-menagerie/zebra_finch |
Bai S, Kolter JZ, Koltun V. An Empirical Evaluation of Generic Convolutional and Recurrent Networks for Sequence Modeling. ArXiv. 2018; abs/1803.01271.

Baker C, Clemens J, Murthy M. Acoustic Pattern Recognition and Courtship Songs: Insights from Insects. Annual Review of Neuroscience. 2019; 42(1):129–147. doi: 10.1146/annurev-neuro-080317-061839.

Bath DE, Stowers JR, Hörmann D, Poehlmann A, Dickson BJ, Straw AD. FlyMAD: rapid thermogenetic control of neuronal activity in freely walking Drosophila. Nature methods. 2014 May; .

Behr O, von Helversen O. Bat serenades—complex courtship songs of the sac-winged bat (Saccopteryx bilineata). Behavioral Ecology and Sociobiology. 2004 Jun; 56(2):106–115.

Benichov JI, Vallentin D. Inhibition within a premotor circuit controls the timing of vocal turn-taking in zebra finches. Nature communications. 2020 Jan; 11(1):1–10.

Bennet-Clark HC. Size and scale effects as constraints in insect sound communication. Philosophical Transactions of the Royal Society B: Biological Sciences. 1998 Mar; 353(1367):407–419.

Calhoun AJ, Pillow JW, Murthy M. Unsupervised identification of the internal states that shape natural behavior. Nature neuroscience. 2019 Nov; 16:1–10.

Campello RJGB, Moulavi D, Sander J. Density-Based Clustering Based on Hierarchical Density Estimates. In: Advances in Knowledge Discovery and Data Mining Berlin, Heidelberg: Springer, Berlin, Heidelberg; 2013.p.160–172.

Cäsar C, Zuberbühler K, Young RJ, Byrne RW. Titi monkey call sequences vary with predator location and type. Biology Letters. 2013 Oct; 9(5):20130535.

Cator LJ, Arthur BJ, Harrington LC, Hoy R. Harmonic Convergence in the Love Songs of the Dengue Vector Mosquito. Science. 2009; 323(5917):1166541+–1166541.

Chaverri G, Gillam EH, Kunz TH. A call-and-response system facilitates group cohesion among disc-winged bats. Behavioral Ecology. 2012 Nov; 24(2):481–487.

Chen X, He K. Exploring Simple Siamese Representation Learning. arXiv:20110566 [cs]. 2020 Nov; .

Choi K, Joo D, Kim J. Kapre: On-GPU Audio Preprocessing Layers for a Quick Implementation of Deep Neural Network Models with Keras. ArXiv. 2017; abs/1706.05781.

Chollet F, et al., Keras; 2015. https://keras.io.

Clay Z, Smith CL, Blumstein DT. Food-associated vocalizations in mammals and birds: what do these calls really mean? Animal Behaviour. 2012 Feb; 83(2):323–330.

Clemens J, Coen P, Roemschied FA, Pereira TD, Mazumder D, Aldarondo DE, Pacheco DA, Murthy M. Discovery of a New Song Mode in Drosophila Reveals Hidden Structure in the Sensory and Neural Drivers of Behavior. Current biology. 2014 Aug; 28(15):2400–2412.e6.

Clemens J, Hennig RM. Computational principles underlying the recognition of acoustic signals in insects. Journal of Computational Neuroscience. 2013 Aug; 35(1):75–85.

Cleveland WS. Robust Locally Weighted Regression and Smoothing Scatterplots. Journal of the American Statistical Association. 1979; .

Coen P, Clemens J, Weinstein AJ, Pacheco DA, Deng Y, Murthy M. Dynamic sensory cues shape song structure in Drosophila. Nature. 2014 Mar; 507(7491):233–237.

Coen P, Xie M, Clemens J, Murthy M. Sensorimotor Transformations Underlying Variability in Song Intensity during Drosophila Courtship. Neuron. 2016 Feb; 89(3):629–644.

Coffey KR, Marx RG, Neumaier JF. DeepSqueak: a deep learning-based system for detection and analysis of ultrasonic vocalizations. Neuropsychopharmacology. 2019 Jan; 231:1–10.

Cohen Y, Nicholson D, Gardner TJ. TweetyNet: A Neural Network That Enables High-Throughput, Automated Annotation of Birdsong. bioRxiv. 2020 Aug; 14(8):2020.08.28.272088. doi: 10.1101/2020.08.28.272088.

Deutsch D, Clemens J, Thiberge SY, Guan G, Murthy M. Shared Song Detector Neurons in Drosophila Male and Female Brains Drive Sex-Specific Behaviors. Current biology. 2019 Oct; 29(19):3200–3215.e5.
Devlin J, Chang MW, Lee K, Toutanova K. BERT: Pre-training of Deep Bidirectional Transformers for Language Understanding. In: NAACL; 2019.

Ding Y, Berrocal A, Morita T, Longden KD, Stern D. Natural courtship song variation caused by an intronic retroelement in an ion channel gene. Nature. 2016; 536:329–332.

Ding Y, Lillvis JL, Cande J, Berman GJ, Arthur BJ, Long X, Xu M, Dickson BJ, Stern DL. Neural Evolution of Context-Dependent Fly Song. Current biology. 2019 Mar; 0(0).

Fitch TW, Neubauer J, Herzel H. Calls out of chaos: the adaptive significance of nonlinear phenomena in mammalian vocal production. Animal Behaviour. 2002; 63(3):407–418.

Fortune ES, Rodríguez C, Li D, Ball GF, Coleman MJ. Neural mechanisms for the coordination of duet singing in wrens. Science. 2011 Nov; 334(6056):666–670.

Gerhardt CH, Huber F. Acoustic Communication in Insects and Anurans. University Of Chicago Press; 2002.

Goffinet J, Brudner S, Mooney R, Pearson J. Low-Dimensional Learned Feature Spaces Quantify Individual and Group Differences in Vocal Repertoires. eLife. 2021 May; 10:e67855. doi: 10.7554/eLife.67855.

Graves A, Jaitly N. Towards End-To-End Speech Recognition with Recurrent Neural Networks. In: International Conference on Machine Learning PMLR; 2014.p. 1764–1772.

Graving JM, Chae D, Naik H, Li L, Koger B, Costelloe BR, Couzin ID. DeepPoseKit, a software toolkit for fast and robust animal pose estimation using deep learning. eLife. 2019 Oct; 8:18.

Guirguis K, Schorn C, Guntoro A, Abdulatif S, Yang B. SELD-TCN: Sound Event Localization & Detection via Temporal Convolutional Networks. 2020 28th European Signal Processing Conference (EUSIPCO). 2021; p. 16–20.

Haack B, Markl H, Ehret G. Sound communication between parents and offspring. In: Willott JF, editor. The auditory psychobiology of the mouse. Springfield (Illinois): C. C. Thomas; 1983.p. 57–97.

Harris CR, Millman KJ, van der Walt SJ, Gommers R, Virtanen P, Cournapeau D, Wieser E, Taylor J, Berg S, Smith NJ, Kern R, Picus M, Hoyer S, van Kerkwijk MH, Brett M, Haldane A, del Rio JF, Wiebe M, Peterson P, Gérard-Marchant P, et al. Array programming with NumPy. Nature. 2020; .

He K, Zhang X, Ren S, Sun J. Deep Residual Learning for Image Recognition. 2016 IEEE Conference on Computer Vision and Pattern Recognition (CVPR). 2016; p. 770–778.

Holy TE, Guo Z. Ultrasonic Songs of Male Mice. PLoS Biology. 2005 Nov; 3(12):e386.

Hoyer S, Hamman J. xarray: N-D labeled arrays and datasets in Python. Journal of Open Research Software. 2017; 5(1). http://doi.org/10.5334/jors.148, doi: 10.5334/jors.148.

Hunter JD. Matplotlib: A 2D graphics environment. Computing in Science & Engineering. 2007; 9(3):90–95. doi: 10.1109/MCSE.2007.55.

Ivanenko A, Watkins P, van Gerven MAJ, Hammerschmidt K, Englitz B. Classifying Sex and Strain from Mouse Ultrasonic Vocalizations Using Deep Learning. PLOS Computational Biology. 2020 Jun; 16(6):e1007918. doi: 10.1371/journal.pcbi.1007918.

Janik VM, Slater PJB. Context-specific use suggests that bottlenose dolphin signature whistles are cohesion calls. Animal Behaviour. 1998 Oct; 56(4):829–838.

Kingma DP, Ba J. Adam: A Method for Stochastic Optimization. CoRR. 2015; abs/1412.6980.

Kluyver T, Ragan-Kelley B, Pérez F, Granger B, Bussonnier M, Frederic J, Kelley K, Hamrick J, Grout J, Corlay S, Ivanov P, Avila D, Abdalla S, Willing C, development team J. Jupyter Notebooks - a publishing format for reproducible computational workflows. In: Loizides F, Schmidt B, editors. Positioning and Power in Academic Publishing: Players, Agents and Agendas Netherlands: IOS Press; 2016. p. 87–90. https://eprints.soton.ac.uk/403913/ .

Kollmorgen S, Hahnloser RHR, Mante V. Nearest neighbours reveal fast and slow components of motor learning. Nature. 2020 Jan; 382:1–5.

Köster J, Rahmann S. Snakemake - a scalable bioinformatics workflow engine. Bioinformatics. 2018; 34 20:3600.
Figure 1–Figure supplement 1. DAS architecture and evaluation.

**A-D** Network architectures for annotating fly song from single (A) and multi-channel (B) recordings, mouse USVs and marmoset song (C), and bird song (Bengalese and Zebra finches) (D). See legend to the right. Each TCN stack consists of stacks of residual blocks shown in E. See Table 4 for all network parameters.

**E** A TCN block (left) consists of a stack of 5 residual blocks (right). Residual blocks process the input with a sequence of dilated convolution, rectification (ReLU) and normalization. The output of this sequence of steps is then added to the input. In successive residual blocks, the dilation rate of the convolution filters doubles from 1x in the first to 16x in the last layer (see numbers to the left of each block). The output of the last residual block is passed as an input to the next TCN block in the network. In addition, the outputs of all residual blocks in a network are linearly combined to predict the song.

**F** Annotation performance is evaluated by comparing manual annotations (top) with labels produced by DAS (bottom). Grey indicates no song, orange song. True negatives (TN) and true positives (TP) are samples for which DAS matches the manual labels. False negatives (FNs) are samples for which the song was missed (blue frame) and reduce recall (TP/(FN+TP)). False positives (FP) correspond to samples that were falsely predicted as containing song (green frames) and reduce precision (TP/(TP+FP)).

**G** Precision and recall are calculated from a confusion matrix which tabulates TP (orange), TN (grey), FP (green), FN (blue). In the example, precision is 3/(3+2) and recall is 3/(1+3).
Figure 1–Figure supplement 2. Performance and the role of context for annotating fly pulse song.

A Recall (blue) and precision (orange) for fly pulse song for different distance thresholds. The distance threshold determines the maximal distance to a true pulse for a detected pulse to be a true positive.

B Waveforms of true positive (left), false positive (middle), and false negatives (right) pulses in fly song. Pulses were aligned to the peak, adjusted for sign, and their amplitude was normalized to have unit norm (see Clemens et al. (2018)).

C Waveforms (top) and confidence scores (bottom, see color bar) for pulses in different contexts. DAS exploits context effects to boost the detection of weak signals. An isolated (“Isolated”, first row) weak pulse-like waveforms is detected with low confidence, since similar waveforms often arise from noise. Manual annotators exploit context information – the fact that pulses often occur in trains at an interval of 40 ms – to annotate weak signals. DAS does the same – the same waveform is detected with much higher confidence due to the presence of a nearby pulse train (“Correct IPI”, 2nd row). If the pulse is too close to another pulse (“Too close”, 3rd row), it is likely noise and DAS detects it with lower confidence. Context effects do not affect strong signals. For instance, a missing pulse within a pulse train (“Missing”, last row) does not reduce detection confidence of nearby pulses.
**Figure 1—Figure supplement 3.** @ plus2 @ minus2 @ plus1 @ plus @ minus2 @

**A** Fly song (black) with manual annotation indicating sine (blue) and pulse (red). Traces (top) and spectrogram (bottom, see color bar) show data from the loudest of the nine audio channels.

**B** Confidence scores (top) for sine (blue) and pulse (red). The confidence is transformed into annotation labels (bottom) based on a confidence threshold (0.5 for sine and pulse). Ground truth (bottom) from manual annotations shown for comparison. DAS annotations were generated using separate networks for pulse or for sine.

**C** Confusion matrix for pulse from a test data set. Color indicates the percentage (see color bar) and text labels indicate number of pulses for each quadrant. All confusion matrices are normalized such that columns sum to 100%. The concentration of values along the diagonal indicates high annotation performance.

**D** Precision-recall curve for pulse depicts the performance characteristics of DAS for different confidence thresholds (from 0 to 1). Recall decreases and precision increases with the threshold. The closer the curve to the upper and right border, the better. The red circle corresponds to the performance of DAS for a threshold of 0.5. The grey circle depicts the performance of FlySongSegmenter (FSS).

**E** Probability density function (PDF) of temporal errors for all detected pulses (red shaded area), computed as the distance between each pulse annotated by DAS and its nearest manual pulse. Lines depict median temporal error for DAS (red line, 0.3 ms) and FSS (grey line, 0.1 ms).

**F, G** Recall of DAS (red line) and FSS (grey line) as a function of the pulse carrier frequency (F) and signal-to-noise ratio (SNR) (G). Red areas show the distributions of carrier frequencies (F) and SNRs (G) for all pulses.

**H** Same as in C but for sine. Color indicates the percentage (see color bar) and text labels indicate seconds of sine for each quadrant.

**I** Same as in D but for sine. The blue circle depicts the performance for the confidence threshold of 0.5 used in A.

**J** Distribution of temporal errors for all detected sine on- and offsets. Median temporal error is 9 ms for DAS (blue line) and 14 ms for FSS (grey line).

**K, L** Recall for DAS (blue line) and FSS (grey line) as a function of sine duration (K) and SNR (L). Blue shaded areas show the distributions of durations (K) and SNRs (L) for all sine songs. DAS outperforms FSS for sine songs with short durations and SNRs <1.0.
Figure 2—Figure supplement 1. Performance for marmoset vocalizations.

A, C Waveform (top) and spectrogram (bottom) of vocalizations from male and female marmosets. Shaded areas (top) show manual annotations of the different vocalization types, colored by type. Recordings are noisy (A, left), clipped (orange), and individual vocalization types are variable (C).

B, D DAS and manual annotation labels for the vocalization types in the recordings in A and C (see color bar in C). DAS annotates the syllable boundaries and types with high accuracy. Note the false negative in D.

E, F Confusion matrices for the four vocalization types in the test set (see color bar), using the syllable boundaries from the manual annotations (E) or from DAS (F) as reference. Rows depict the probability with which DAS annotated each syllable as any of the four types in the test dataset. The type of most syllables were correctly annotated, resulting in the concentration of probability mass along the main diagonal. False positives and false negatives correspond to the first row in E and the first column in F, respectively. When using the true syllables for reference, there are no false positives (E, x="noise", grey labels) since all detections are positives. By contrast, when using the predicted syllables as reference, there are no true negatives (F, y="noise", grey labels), since all reference syllables are (true or false) positives.

G Distribution of temporal errors for the on- and offsets of all detected syllables (purple shaded area). The median temporal error is 4.4 ms for DAS (purple line) and 12.5 ms for the method by Oikarinen et al. (2019) developed to annotate marmoset calls (grey line).
Figure 3–Figure supplement 1. Performance for the song of Bengalese finches. 

A, B Number of syllables (log scaled) in the train (A) and the test (B) sets for each syllable type present in the test set.

C Precision (blue) and recall (orange) for each syllable type, computed from the confusion matrix in Figure 3C.

D Confusion matrices when using true (left, same as Figure 3C) and predicted (right) syllables as a reference. Colors were log-scaled to make the rare annotation errors more apparent (see color bar). The reference determines the syllable bounds and the syllable label is then given by the most frequent label found in the samples within the syllable bounds. When using the true syllables for reference, there are no false positives (left, y=0 (no song), grey line) since all detections are positives. By contrast, when using the predicted syllables as reference, there are no true negatives (right, x=0 (no song), grey line), since all reference syllables are (true or false) positives. The average false negative and positive rates are 0.3% and 0.2%, respectively.
Figure 3–Figure supplement 2. Performance for the song of a Zebra finch.

**A** Waveform (top) and spectrogram (bottom) of the song of a male Zebra finch. Shaded areas (top) show manual annotations of the six syllables of the male’s motif, colored by syllable type.

**B** DAS and manual annotation labels for the six syllable types in the recording in A (see color bar). DAS accurately annotates the syllable boundaries and types.

**C** Confusion matrix for the six syllables in the test set (see color bar). Rows depict the probability with which DAS annotated each syllable as any of the six types in the test dataset. The type of 100% (54/54) of the syllables were correctly annotated, resulting in the concentration of probability mass along the main diagonal.

**D** Distribution of temporal errors for the on- and offsets of all detected syllables in the test set (blue shaded area). The median temporal error is 1.3 ms for DAS (blue line) and 2 ms for TweetyNet (Cohen et al., 2020), a method developed to annotate bird song (grey line).
Figure 4–Figure supplement 1. Throughput and latency of inference.
Median throughput (A-H) and latency (I-P) across 10 inference runs for different song types. Batch duration is the product of chunk duration and batch size. Error bars are smaller than the marker size (see Figure 4A/B). Solid and dashed lines correspond to values when running DAS on a CPU and GPU, respectively. Squares and circles in the plots for fly song correspond to networks with short and long chunk durations, respectively (see Figure 4–Figure Supplement 2). In A-H, horizontal lines mark 10x and 100x realtime and vertical lines mark the batch duration used in Figure 4A. In I-P, the vertical line marks the batch duration used in Figure 4B. Differences in the throughput and latency between species arise from differences in the sample rate and the complexity of the network (number and duration of the filters, number of TCN blocks) (see Table 4).
Figure 4-Figure supplement 2. Reducing the chunk duration reduces latency and comes with minimal performance penalties.

A F1 scores of networks with different chunk durations. Reduction of performance due to loss of context information with shorter chunks is minimal.

B Latency of networks with different chunk durations. Dashed vertical lines indicate the chunk durations of the networks in Figure 1, Figure 4B and Figure 4-Figure Supplement 1. Solid vertical lines indicated the “short” chunk durations used in Figure 4B and Figure 4-Figure Supplement 1.
**Figure 4—Figure supplement 3. DAS requires small to moderate amounts of data for training.**

A Performance as a function of the number of manual annotations in the training set. Individual dots correspond to individual fits using different subsets of the data. For flies and mice, dots indicate the number of annotations for each type and performance is given by the F1 score—the geometric mean of precision and recall. For the marmosets and birds (rightmost panels), dots correspond to the median number of syllables per type in the training set and performance is given by the median accuracy over syllables per fit (see C for per-type statistics for Bengalese finches). Colored curves in A and B were obtained by locally weighted scatter plot smoothing (lowess) of the individual data points. Vertical lines correspond to the number of syllables required to surpass 90% of the maximal performance.

B Temporal error given as the median temporal distance to syllable on- and offsets or to pulses.

C Number of annotations required per syllable type for Bengalese finches (range 2-64, median 8, mean 17). One outlier type requires 200 annotations and consists of a mixture of different syllable types (black bar, see D-F).

D, E UMAP embedding of the bird syllables with the outlier type from C labelled according to the manual annotations (D, black) and the unsupervised clustering (F). The unsupervised clustering reveals that the outlier type splits into two distinct clusters of syllables (cyan and pink) and three mislabelled syllables (red).

F Spectrograms of different types of syllable exemplars from the outlier type in C grouped based on the unsupervised clustering in E.

G Confusion matrix mapping manual to unsupervised cluster labels (see Figure 5F, G) for marmosets. Green boxes mark the most likely call type for each unsupervised cluster. While there is a good correspondence between manual and unsupervised call types, most call types are split into multiple clusters.
DAS can be trained using an iterative procedure, in which the network is network trained after annotating few syllables. This initial network is then used to create a larger dataset of annotations, by predicting and manually correcting annotation labels for a longer stretch of audio. This procedure is repeated to create ever larger training datasets until the performance is satisfactory. Shown is an example from mouse song.

A Example of the test recording for mouse song (top - waveform, bottom - spectrogram)
B Manual labels (true, black) and correct (blue) and erroneous (red) annotations from DAS for three different stages of training. Even for the first round of fast training, the majority of onsets is detected with low temporal error, requiring only few corrections. Number of syllables, seconds of song and F1-score for the different iterations (1st/2nd/final): 72/248/2706 syllables, 6/26/433 seconds of song, F1 score 96.6/97.9/98.9.
**Figure 4–Figure supplement 5.** DAS performance is robust to changes in the structural parameters of the network.

A, B Performance (F1 score) for networks with different structural parameters for fly pulse (A, red) and sine (B, blue). The starting point for each modified network was the network given in Table 4. We then trained network with individual modified parameters from scratch. As long as the network has chunks longer than 100 ms, filters longer than 2 ms, more than 8 filters, and at least one TCN block, the resulting model has an F1 score of 95% for pulse and sine.

C Same in B but for Bengalese finches. Network performance is robust to changes in individual parameters. Convergence is inconsistent for shallow networks with 1-2 TCN blocks (rightmost panel).

F1 scores smaller than 80% in A-C where set to 80% to highlight small changes in performance.