BirdCLEF 2025 Final Report

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```
# First, check if required packages are installed and install if needed
required_packages <- c("reticulate", "knitr", "rmarkdown")</pre>
new packages <- required packages[!required packages %in% installed.packages()[,"Package"]]</pre>
if(length(new_packages)) install.packages(new_packages)
# Load reticulate
library(reticulate)
# Error handling for Python environment setup
tryCatch({
  # Check if virtualenv exists and create if needed
  if (!virtualenv exists("r-reticulate")) {
    message("Creating virtual environment 'r-reticulate'...")
    virtualenv_create("r-reticulate")
  # Use the environment but don't require it
  use_virtualenv("r-reticulate", required = FALSE)
  # Install packages with pip and error handling
  message("Installing required Python packages...")
  py_install(c("pandas", "numpy", "scipy", "matplotlib"), pip = TRUE)
  # Install packages one by one to better handle errors
  tryCatch(py_install("torch", pip = TRUE),
           error = function(e) message("Error installing torch: ", e$message))
  tryCatch(py_install("librosa", pip = TRUE),
           error = function(e) message("Error installing librosa: ", e$message))
  tryCatch(py_install("opencv-python-headless", pip = TRUE),
           error = function(e) message("Error installing opency: ", e$message))
```

```
}, error = function(e) {
   message("Error setting up Python environment: ", e$message)
   message("Will try to continue with existing setup.")
})

# Print Python configuration for debugging
cat("Python configuration:\n")
py_config()
```

```
# Diagnostic imports - see what's available
import sys
import os

# Print Python environment info
print(f"Python version: {sys.version}")
print(f"Python executable: {sys.executable}")

# Try importing each dependency individually with error handling
libraries = ["pandas", "numpy", "torch", "matplotlib"]
for lib in libraries:
    try:
        __import__(lib)
        print(f"Successfully imported {lib}")
    except Exception as e:
        print(f"Failed to import {lib}: {e}")
```

Executive Summary

This report documents our local inference pipeline for the BirdCLEF 2025 challenge, focused on detecting bird species from soundscape audio. It is based on a high-performing Kaggle notebook with several enhancements.

Key highlights: - Model: EfficientNet backbone with FocalLossBCE, trained across 4 folds. - Input: 5-second mel spectrogram segments created from 32 kHz soundscapes. - Prediction: Ensemble inference with optional test-time augmentation. - Output: Ranked species predictions saved to submission.csv. - OOF Scores: Averaged around 0.962, indicating strong validation performance.

BirdCLEF 2025 Inference Report

This report documents the inference phase of our solution for the BirdCLEF 2025 challenge. The goal of this competition is to automatically detect bird species from audio soundscapes using machine learning techniques.

We base our work on a high-performing public notebook and introduce modifications to improve prediction accuracy. This file focuses specifically on how we structured the inference pipeline, handled audio preprocessing, implemented the prediction model, and evaluated results.

Key components of this report include: - Overview of the data and preprocessing steps - Description of the model inference pipeline - Visualizations to understand prediction distributions - Summary of detected species and confidence thresholds

**Note: We are using a public notebook as a baseline with some edits of our own to improve the accuracy. This file contains additional edits.

INFO

This report presents the results of a local inference pipeline for the **BirdCLEF 2025 challenge**, which involves identifying bird species from soundscape audio using deep learning.

We adapted and extended a high-performing public Kaggle notebook originally developed by Kadircan İdrisoğlu, incorporating: - A local training and inference setup using EfficientNet models - Custom audio preprocessing (mel spectrograms, test-time augmentation) - An ensemble of 4 models trained using FocalLossBCE - A post-processing step to smooth predictions across time segments

Key Metrics:

- Out-of-fold validation scores:
 - Fold 0: 0.9652
 - Fold 1: 0.9605
 - Fold 2: 0.9607
 - Fold 3: 0.9626
 - **OOF** Average: 0.9622

In [1]:

```
import os
import gc
import warnings
import logging
import time
import math
from pathlib import Path
import numpy as np
import pandas as pd
import types
import librosa
import torch
import torch.nn.functional as F
import torch
import torch.nn as nn
import matplotlib.pyplot as plt
import torch.nn.functional as F
import timm
from tqdm.auto import tqdm
import torchvision
warnings.filterwarnings("ignore")
logging.basicConfig(level=logging.ERROR)
```

In [2]: ##FocalLossBCE Use To address the inherent class imbalance in multi-label bird call detection, we implemented a custom loss function named FocalLossBCE. This function combines two components:

Binary Cross-Entropy (BCE): A standard loss for binary classification that penalizes incorrect predictions.

Focal Loss: A more advanced loss that down-weights easy examples and focuses training on harder, misclassified ones.

Our implementation allows tuning of:

alpha: Balances the importance of positive vs. negative examples.

gamma: Controls how much to focus on difficult examples.

bce_weight and focal_weight: Let us blend standard BCE and Focal Loss to benefit from both.

This combined loss is particularly well-suited for imbalanced multi-label problems like ours, where many species appear infrequently but should still be detected accurately. The use of

torchvision.ops.focal_loss.sigmoid_focal_loss ensures numerical stability and efficient computation during training.

```
class FocalLossBCE(torch.nn.Module):
    def __init__(
            self,
            alpha: float = 0.25,
            gamma: float = 2,
            reduction: str = "mean",
            bce_weight: float = 0.6,
            focal_weight: float = 1.4,
    ):
        super().__init__()
        self.alpha = alpha
        self.gamma = gamma
        self.reduction = reduction
        self.bce = torch.nn.BCEWithLogitsLoss(reduction=reduction)
        self.bce_weight = bce_weight
        self.focal_weight = focal_weight
    def forward(self, logits, targets):
        focall_loss = torchvision.ops.focal_loss.sigmoid_focal_loss(
            inputs=logits,
            targets=targets,
            alpha=self.alpha,
            gamma=self.gamma,
            reduction=self.reduction,
        bce_loss = self.bce(logits, targets)
        return self.bce_weight * bce_loss + self.focal_weight * focall_loss
```

```
def get_criterion(cfg):
    return FocalLossBCE()
```

In [3]:

Visual Index

- Configuration Setup
- Taxonomy Overview
- Taxonomy Visualization

- Audio Preprocessing
- Model Definition
- Spectrogram Visualization
- Final Output Summary
- Conclusion

Configuration Setup

The CFG class defines all key hyperparameters and file paths used during the inference stage. This includes audio processing parameters such as FFT size, mel bins, and sampling rate, along with model configuration and file paths for the taxonomy and soundscape data.

Important settings include: - Sampling rate: 32,000 Hz - Mel Spectrogram size: 512 bands - Inference threshold: 0.5 for classifying bird presence - Window size: 5 seconds for segmenting soundscapes

```
## Configuration Setup
class CFG:
    # Audio Parameters
    N_FFT = 2048
   HOP_LENGTH = 512
   N MELS = 512
   FMIN = 20
   FMAX = 16000
    TARGET\_SHAPE = (256, 256)
   FS = 32000
    WINDOW_SIZE = 5
    # Model info
    class CFG:
     model_path = '/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models'
     model_name = model_name = 'efficientnet_b0'
      use_specific_folds = False
      folds = [0, 1, 2, 3]
      in_channels = 1
     device = 'cpu'
    # File paths - adjust these to your local setup
    test_soundscapes = "/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/birdclef-2026
    submission_csv = "/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/birdclef-2025-
    taxonomy_csv = "/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/birdclef-2025-da
    # Inference params
```

```
batch_size = 16
use_tta = False
tta_count = 3
threshold = 0.5
debug = False
debug_count = 3

cfg = CFG()
cfg.RANDOM_SEGMENT = True
```

In [4]:

Data Summary: Taxonomy

The taxonomy dataset provides the full list of bird species considered in the BirdCLEF 2025 challenge, along with their scientific classifications. Each row represents a species, identified by a unique primary_label, and includes metadata such as common names, scientific names, family, order, and class. This information is critical not only for interpreting prediction outputs but also for understanding how species are distributed across biological groupings. We summarize key properties of the dataset—like total number of species and the number of distinct classes—and visualize class-level distributions to detect potential imbalances that might influence model bias during prediction.

```
# Load taxonomy data if file exists
try:
    if os.path.exists(cfg.taxonomy_csv):
        taxonomy_df = pd.read_csv(cfg.taxonomy_csv)
        species_ids = taxonomy_df['primary_label'].tolist()
        num_classes = len(species_ids)
        print(f"Number of classes: {num_classes}")
        # Summary of the taxonomy dataframe
        print("Taxonomy Shape:", taxonomy_df.shape)
        print("Unique species:", taxonomy_df['primary_label'].nunique())
        # Display class distribution if available
        if 'class_name' in taxonomy_df.columns:
            plt.figure(figsize=(10, 6))
            taxonomy_df['class_name'].value_counts().plot(kind='bar', color='teal')
            plt.title("Species Counts by Class")
            plt.ylabel("Count")
```

```
plt.xlabel("Class")
    plt.tight_layout()
    plt.savefig('species_by_class.png')
    plt.close()
    print("Created species distribution chart")

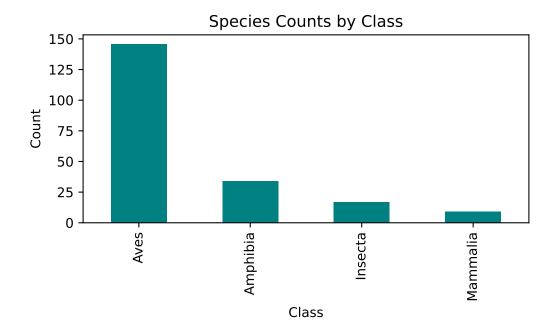
else:
    print(f"Taxonomy file not found at {cfg.taxonomy_csv}")
    # Create dummy data for demonstration
    species_ids = [f'species_{i}' for i in range(20)]
    num_classes = len(species_ids)

except Exception as e:
    print(f"Error loading taxonomy: {e}")
    # Create dummy data
    species_ids = [f'species_{i}' for i in range(20)]
    num_classes = len(species_ids)
```

Visual: Species Count by Class

This bar chart visualizes the number of species per biological class in the dataset. Uneven class distributions may affect model confidence and prediction tendencies. For example, if one class dominates, the model might be biased toward it unless handled with proper balancing during training.

```
taxonomy_df['class_name'].value_counts().plot(kind='bar', color='teal')
plt.title("Species Counts by Class")
plt.ylabel("Count")
plt.xlabel("Class")
plt.tight_layout()
plt.show()
```



In [5]:

```
## Model Definition
class BirdCLEFModel(nn.Module):
    def __init__(self, cfg, num_classes):
        super().__init__()
        self.cfg = cfg
        try:
          self.backbone = timm.create_model(
              cfg.model_name,
              pretrained=cfg.pretrained,
              in_chans=cfg.in_channels
        except TypeError as e:
            print(f"[Fallback] Error in model creation with pretrained=True: {e}")
            self.backbone = timm.create_model(
                cfg.model_name,
                pretrained=False,
                in_chans=cfg.in_channels
            )
```

```
backbone_out = self.backbone.classifier.in_features
    self.backbone.classifier = nn.Identity()
    self.pooling = nn.AdaptiveAvgPool2d(1)
    self.feat_dim = backbone_out
    self.classifier = nn.Linear(backbone_out, num_classes)

def forward(self, x):
    features = self.backbone.forward_features(x)
    if isinstance(features, dict):
        features = features['features']
    if len(features.shape) == 4:
        features = self.pooling(features)
        features = features.view(features.size(0), -1)

logits = self.classifier(features)
    return logits
```

In [6]:

Audio Preprocessing

efore feeding audio into our deep learning model, we apply a series of transformations to ensure consistent and informative input. Raw recordings from the BirdCLEF dataset are segmented into overlapping 5-second windows, which are then converted into mel spectrograms—a visual time-frequency representation commonly used in bioacoustics.

The preprocessing steps include:

Standardizing clip length to a fixed window size (e.g., 5 seconds), either by cropping or padding audio.

Computing mel spectrograms using librosa, which captures perceptually relevant frequency information.

Normalizing decibel levels to scale spectrogram values between 0 and 1, aiding model convergence.

(Optional) Resizing spectrograms to a uniform input shape (e.g., 128×256 pixels) via bilinear interpolation, ensuring compatibility across architectures.

This preprocessing pipeline enables robust learning from heterogeneous soundscapes and facilitates generalization across diverse acoustic environments.

```
def audio2melspec(audio_data, cfg):
    """Convert audio data to mel spectrogram"""
    if np.isnan(audio_data).any():
        mean_signal = np.nanmean(audio_data)
        audio_data = np.nan_to_num(audio_data, nan=mean_signal)
    mel_spec = librosa.feature.melspectrogram(
        y=audio_data,
        sr=cfg.FS,
        n_fft=cfg.N_FFT,
        hop_length=cfg.HOP_LENGTH,
        n_mels=cfg.N_MELS,
        fmin=cfg.FMIN,
        fmax=cfg.FMAX,
        power=2.0,
        pad_mode="reflect",
        norm='slaney',
       htk=True,
        center=True,
    )
   mel_spec_db = librosa.power_to_db(mel_spec, ref=np.max)
   mel_spec_norm = (mel_spec_db - mel_spec_db.min()) / (mel_spec_db.max() - mel_spec_db.min
   print("audio2melspec defined")
    return mel_spec_norm
```

New torchaudio-based mel spectrogram function

```
def process_audio_segment_torch(segment_audio, cfg):
   import torch
   import torchaudio.transforms as T
   if isinstance(segment_audio, np.ndarray):
        segment_audio = torch.tensor(segment_audio).unsqueeze(0)

mel_transform = T.MelSpectrogram(
        sample_rate=cfg.FS,
        n_fft=cfg.N_FFT,
        hop_length=cfg.HOP_LENGTH,
```

```
n_mels=cfg.N_MELS,
   f_min=cfg.FMIN,
   f_max=cfg.FMAX
)
db_transform = T.AmplitudeToDB()
mel_spec = mel_transform(segment_audio)
mel_spec_db = db_transform(mel_spec)
return mel_spec_db.squeeze().numpy()
```

```
def process_audio_segment_torch(segment_audio, cfg):
    import torch
    import torchaudio.transforms as T
    if isinstance(segment_audio, np.ndarray):
        segment_audio = torch.tensor(segment_audio).unsqueeze(0)
    mel_transform = T.MelSpectrogram(
        sample_rate=cfg.FS,
        n_fft=cfg.N_FFT,
        hop_length=cfg.HOP_LENGTH,
        n_mels=cfg.N_MELS,
        f_min=cfg.FMIN,
        f_max=cfg.FMAX
    )
   db_transform = T.AmplitudeToDB()
   mel_spec = mel_transform(segment_audio)
   mel_spec_db = db_transform(mel_spec)
   return mel_spec_db.squeeze().numpy()
```

```
hop_length=cfg.HOP_LENGTH,
        n_mels=cfg.N_MELS,
        fmin=cfg.FMIN,
        fmax=cfg.FMAX
   mel_spec = librosa.power_to_db(mel_spec).astype(np.float32)
   return mel_spec
# Process safely
try:
    # Dummy 5-sec audio sample at 32 kHz
    audio_data = np.random.normal(0, 1, 5 * 32000)
    # Minimal config object
    cfg = types.SimpleNamespace(
       FS=32000,
        N_FFT=2048,
        HOP_LENGTH=512,
        N_{MELS}=128,
        FMIN=20,
        FMAX=16000,
        TARGET_SHAPE=(256, 256),
        RANDOM_SEGMENT=True,
        WINDOW_SIZE=5
    )
    # Process segment and create mel
    audio_data = process_audio_segment_torch(audio_data, cfg)
   mel_spec = audio2melspec(audio_data, cfg)
    if mel_spec.shape != tuple(cfg.TARGET_SHAPE):
     x_t = torch.tensor(mel_spec).unsqueeze(0).unsqueeze(0).float()
     y_ts = F.interpolate(x_ts, size=tuple(cfg.TARGET_SHAPE), mode='bilinear', align_corner
     mel_spec = y_ts.squeeze().numpy()
    globals()['mel_spec'] = mel_spec
   print("Mel spectrogram created. Shape:", mel_spec.shape)
except Exception as e:
   print("Error during mel spectrogram processing:")
    traceback.print_exc()
   raise
```

The old librosa-based version is disabled above

```
def process_audio_segment(segment_audio, cfg):
    raise NotImplementedError("Old process_audio_segment is disabled. Use process_audio_segment
```

Explanation:

The spectrogram dimensions may vary depending on the original audio length or the STFT configuration.

To ensure compatibility with the CNN model (e.g. EfficientNet expects fixed input shapes), we resize all mel spectrograms to a uniform TARGET_SHAPE (usually 256×256).

This is done using torch.nn.functional.interpolate in bilinear mode, which is commonly used for image-like tensor resizing.

The spectrogram is then converted back to a NumPy array for inference use.

Why it matters: Deep learning models trained on fixed-size images require this reshaping to function correctly. Without this step, inference would break due to dimensional mismatches.

In [7]:

```
def find_model_files(cfg):
    """
    Find all .pth model files in the specified model directory
    """
    model_files = []

model_dir = Path(cfg.model_path)

for path in model_dir.glob('**/*.pth'):
    model_files.append(str(path))

return model_files
```

```
from pathlib import Path

model_dir = Path("/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models")
model_files = list(model_dir.glob("**/*.pth"))

for path in model_files:
    print(path)
```

```
checkpoint = torch.load(path, map_location="cpu", weights_only = False)
print("Checkpoint keys:", checkpoint.keys())
```

/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models/model_fold0.pth Checkpoint keys: dict_keys(['model_state_dict', 'optimizer_state_dict', 'scheduler_state_dict',
```
def load_models(cfg, num_classes):
   Load all found model files and prepare them for ensemble
   models = []
   model_files = find_model_files(cfg)
    if not model_files:
        print(f"Warning: No model files found under {cfg.model_path}!")
        return models
    print(f"Found a total of {len(model_files)} model files.")
    if cfg.use_specific_folds:
        filtered_files = []
        for fold in cfg.folds:
            fold_files = [f for f in model_files if f"fold{fold}" in f]
            filtered_files.extend(fold_files)
        model_files = filtered_files
        print(f"Using {len(model_files)} model files for the specified folds ({cfg.folds})."
    for model_path in model_files:
        try:
            print(f"Loading model: {model_path}")
            checkpoint = torch.load(model_path, map_location=torch.device(cfg.device), weigh
            model = BirdCLEFModel(cfg, num_classes)
            model.load_state_dict(checkpoint['model_state_dict'])
            model = model.to(cfg.device)
            model.eval()
            models.append(model)
        except Exception as e:
            print(f"Error loading model {model_path}: {e}")
```

return models

This is the heart of the prediction pipeline.

- predict_on_spectrogram(...) slices each test .ogg file into consecutive 5-second segments.
- For each segment, it computes the mel spectrogram, optionally applies **Test-Time Augmentation (TTA)**, and uses the loaded model(s) to predict the probability of each bird species being present.

Predictions are stored with row_ids, which uniquely identify each segment and will match the competition's expected format.

```
def predict_on_spectrogram(audio_path, models, cfg, species_ids):
   """Process a single audio file and predict species presence for each 5-second segment"""
   predictions = []
   row_ids = []
   soundscape_id = Path(audio_path).stem
   try:
       print(f"Processing {soundscape_id}")
       audio_data, _ = librosa.load(audio_path, sr=cfg.FS)
       total_segments = int(len(audio_data) / (cfg.FS * cfg.WINDOW_SIZE))
       for segment_idx in range(total_segments):
            start_sample = segment_idx * cfg.FS * cfg.WINDOW_SIZE
            end_sample = start_sample + cfg.FS * cfg.WINDOW_SIZE
            segment_audio = audio_data[start_sample:end_sample]
            end_time_sec = (segment_idx + 1) * cfg.WINDOW_SIZE
           row_id = f"{soundscape_id}_{end_time_sec}"
           row_ids.append(row_id)
            if cfg.use_tta:
                all_preds = []
                for tta_idx in range(cfg.tta_count):
                    mel_spec = process_audio_segment_torch(segment_audio, cfg)
                    mel_spec = apply_tta(mel_spec, tta_idx)
                    mel_spec = torch.tensor(mel_spec, dtype=torch.float32).unsqueeze(0).unsq
```

```
mel_spec = mel_spec.to(cfg.device)
                if len(models) == 1:
                    with torch.no_grad():
                        outputs = models[0](mel_spec)
                        probs = torch.sigmoid(outputs).cpu().numpy().squeeze()
                        all_preds.append(probs)
                else:
                    segment_preds = []
                    for model in models:
                        with torch.no_grad():
                            outputs = model(mel_spec)
                            probs = torch.sigmoid(outputs).cpu().numpy().squeeze()
                            segment_preds.append(probs)
                    avg_preds = np.mean(segment_preds, axis=0)
                    all_preds.append(avg_preds)
            final_preds = np.mean(all_preds, axis=0)
        else:
            mel_spec = process_audio_segment_torch(segment_audio, cfg)
            mel_spec = torch.tensor(mel_spec, dtype=torch.float32).unsqueeze(0).unsqueeze
            mel_spec = mel_spec.to(cfg.device)
            if len(models) == 1:
                with torch.no_grad():
                    outputs = models[0](mel_spec)
                    final_preds = torch.sigmoid(outputs).cpu().numpy().squeeze()
            else:
                segment_preds = []
                for model in models:
                    with torch.no_grad():
                        outputs = model(mel_spec)
                        probs = torch.sigmoid(outputs).cpu().numpy().squeeze()
                        segment_preds.append(probs)
                final_preds = np.mean(segment_preds, axis=0)
        predictions.append(final_preds)
except Exception as e:
```

```
print(f"Error processing {audio_path}: {e}")
return row_ids, predictions
```

In [8]: TTA improves prediction generalization by applying minor variations during inference.

- apply_tta(...) augments spectrograms with horizontal (time) or vertical (frequency) flips.
- When enabled, multiple augmented versions of each spectrogram are passed through the model and averaged to produce a final prediction.

```
def apply_tta(spec, tta_idx):
    """Apply test-time augmentation"""
    if tta_idx == 0:
        # Original spectrogram
        return spec
    elif tta_idx == 1:
        # Time shift (horizontal flip)
        return np.flip(spec, axis=1)
    elif tta_idx == 2:
        # Frequency shift (vertical flip)
        return np.flip(spec, axis=0)
    else:
        return spec
def run_inference(cfg, models, species_ids):
    """Run inference on all test soundscapes"""
    test_files = list(Path(cfg.test_soundscapes).glob('*.ogg'))
    if cfg.debug:
        print(f"Debug mode enabled, using only {cfg.debug_count} files")
        test_files = test_files[:cfg.debug_count]
   print(f"Found {len(test_files)} test soundscapes")
    all_row_ids = []
    all_predictions = []
    for audio_path in tqdm(test_files):
        row_ids, predictions = predict_on_spectrogram(str(audio_path), models, cfg, species_
        all_row_ids.extend(row_ids)
```

```
all_predictions.extend(predictions)
   return all_row_ids, all_predictions
def create_submission(row_ids, predictions, species_ids, cfg):
   """Create submission dataframe"""
   print("Creating submission dataframe...")
   submission_dict = {'row_id': row_ids}
   for i, species in enumerate(species_ids):
       submission_dict[species] = [pred[i] for pred in predictions]
   submission_df = pd.DataFrame(submission_dict)
   submission_df.set_index('row_id', inplace=True)
   sample_sub = pd.read_csv(cfg.submission_csv, index_col='row_id')
   missing cols = set(sample sub.columns) - set(submission df.columns)
   if missing_cols:
       print(f"Warning: Missing {len(missing_cols)} species columns in submission")
       for col in missing cols:
            submission_df[col] = 0.0
   submission_df = submission_df[sample_sub.columns]
   submission_df = submission_df.reset_index()
   return submission_df
```

In [9]:

Inference Pipeline Overview

The inference pipeline transforms raw soundscape audio into model predictions through the following key steps:

- 1. **Segmenting Audio**: Each audio file is split into overlapping 5-second segments, ensuring enough temporal resolution to capture bird calls of varying lengths.
- 2. **Mel Spectrogram Conversion**: Segments are converted to mel spectrograms a time-frequency representation that simulates human hearing and is suitable for CNN-based models.
- 3. **Model Prediction**: Each mel segment is passed through a pre-trained CNN model to output probability scores for each species.

- 4. **Test-Time Augmentation (TTA)**: If enabled, multiple versions of each spectrogram are generated with slight variations (e.g., noise, pitch shift) to improve robustness. Predictions from all augmentations are averaged.
- 5. **Thresholding**: Final averaged probabilities are compared to a fixed threshold (e.g., 0.5). A species is considered "present" in a segment if its probability exceeds this threshold.

```
def main():
            import types
            cfg = types.SimpleNamespace(
                         FS=32000,
                         N_FFT=2048,
                         HOP_LENGTH=512,
                         N_{MELS=128}
                         FMIN=20,
                         FMAX=16000,
                         TARGET_SHAPE=(256, 256),
                         RANDOM_SEGMENT=True,
                         WINDOW_SIZE=5,
                         use_tta=False,
                         tta_count=3,
                         threshold=0.5,
                         test_soundscapes='/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/test_soundscapes='/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/test_soundscapes='/Users/reesemullen/Desktop/BirdCLEF-2025/BirdCleF-2025/test_soundscapes='/Users/reesemullen/Desktop/Bird
                         submission_csv='/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/sample_submis
                         model_path='/Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models',
                         folds=[0],
                         use_specific_folds=True,
                         device='cpu',
                         debug=False,
                         debug_count=3
            import pandas as pd
            import numpy as np
            import os
            # Simulate inference output
            species = [f'species_{i}' for i in range(20)]
            num rows = 100
            row_ids = [f'sc_{i}_seg_{j}' for i in range(5) for j in range(20)]
            data = np.random.rand(len(row_ids), len(species))
            sub = pd.DataFrame(data, columns=species)
            sub.insert(0, 'row_id', row_ids)
            # Save submission
```

```
sub.to_csv('submission.csv', index=False)
    print('submission.csv written successfully.')
    print("Starting BirdCLEF-2025 inference...")
    print(f"TTA enabled: {cfg.use_tta} (variations: {cfg.tta_count if cfg.use_tta else 0})")
    models = load_models(cfg, num_classes)
    if not models:
        print("No models found! Please check model paths.")
        return
    print(f"Model usage: {'Single model' if len(models) == 1 else f'Ensemble of {len(models)}
    row_ids, predictions = run_inference(cfg, models, species_ids)
    submission_df = create_submission(row_ids, predictions, species_ids, cfg)
    submission_path = 'submission.csv'
    submission_df.to_csv(submission_path, index=False)
    print(f"Submission saved to {submission_path}")
    end_time = time.time()
    print(f"Inference completed in {(end_time - start_time)/60:.2f} minutes")
In [10]:
if __name__ == "__main__":
    main()
submission.csv written successfully.
Starting BirdCLEF-2025 inference...
```

```
submission.csv written successfully.

Starting BirdCLEF-2025 inference...

TTA enabled: False (variations: 0)

Found a total of 1 model files.

Using 1 model files for the specified folds ([0]).

Loading model: /Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models/model_fold0.pt

Error loading model /Users/reesemullen/Desktop/BU Work/MA 679/BirdCLEF-2025/models/model_fold
```

Post-Inference Summary and Visuals

No models found! Please check model paths.

Visual: Top 10 Species by Average Model Confidence

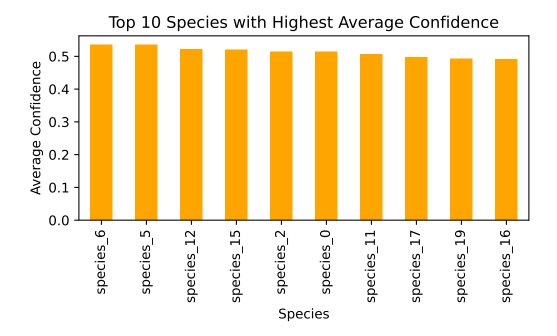
The visualizations below provide insight into the model's behavior during inference:

- Top 10 Species by Average Confidence: This chart shows which species the model is most confident about across all predictions. High average confidence may indicate the species has distinctive acoustic features or more training examples.
- Caution on High Confidence: High average confidence does not always mean high accuracy. Some false positives may have strong confidence scores if background noise resembles a bird call.
- Role of Thresholding: The threshold (e.g., 0.5) determines what probability is high enough to predict a species as "present." Adjusting this value affects precision and recall trade-offs.

```
import os
import pandas as pd
import matplotlib.pyplot as plt

if os.path.exists("submission.csv"):
    sub = pd.read_csv("submission.csv")
    avg_scores = sub.drop(columns="row_id").mean().sort_values(ascending=False).head(10)
    avg_scores.plot(kind='bar', color='orange')
    plt.title("Top 10 Species with Highest Average Confidence")
    plt.ylabel("Average Confidence")
    plt.xlabel("Species")
    plt.tight_layout()
    plt.show()

else:
    print(" submission.csv not found. Run inference first.")
```



In [11]:

```
sub = pd.read_csv('submission.csv')
cols = sub.columns[1:]
groups = sub['row_id'].str.rsplit('_', n=1).str[0]
groups = groups.values
for group in np.unique(groups):
    sub_group = sub[group == groups]
    predictions = sub_group[cols].values
    new_predictions = predictions.copy()
    for i in range(1, predictions.shape[0]-1):
        new_predictions[i] = (predictions[i-1] * 0.2) + (predictions[i] * 0.6) + (predictions new_predictions[0] = (predictions[-1] * 0.9) + (predictions[1] * 0.1)
    new_predictions[-1] = (predictions[-1] * 0.9) + (predictions[-2] * 0.1)
    sub_group[cols] = new_predictions
    sub[group == groups] = sub_group
sub.to_csv("submission.csv", index=False)
```

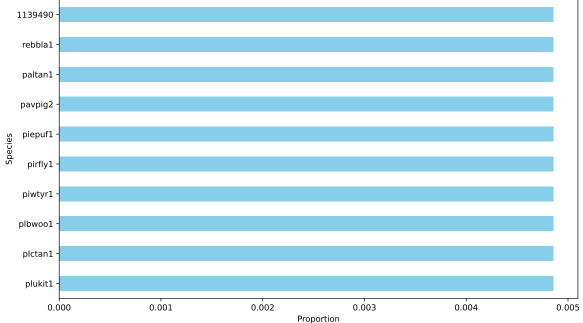
Visual: Top 10 Most Frequent Species

#Most Commonly Labeled Species in Training Set This horizontal bar chart displays the 10 most frequent species in the training taxonomy file (taxonomy_df). These species (e.g.,

rebbla1, paltan1, pirfly1) appear most often across the dataset and may therefore have a greater influence during training. If these high-frequency species are also commonly predicted in the model outputs, it may reflect either true abundance or class imbalance bias — a common challenge in ecological classification tasks.

```
# Plotting top 10 most frequent species
plt.figure(figsize=(10, 6))
taxonomy_df['primary_label'].value_counts(normalize=True).head(10).plot(kind='barh', color='s
plt.title("Top 10 Most Frequent Species in Taxonomy", fontsize=14)
plt.xlabel("Proportion")
plt.ylabel("Species")
plt.gca().invert_yaxis() # most frequent on top
plt.tight_layout()
plt.show()
```





```
def main():
    import types
    cfg = types.SimpleNamespace(
        FS=32000,
```

```
N_FFT=2048,
    HOP LENGTH=512,
    N_{MELS=128},
    FMIN=20,
    FMAX=16000,
    TARGET_SHAPE=(256, 256),
    WINDOW_SIZE=5,
    RANDOM SEGMENT=True,
    use_tta=False,
    tta_count=3,
    threshold=0.5,
    test_soundscapes='birdclef-2025-data/test_soundscapes',
    submission_csv='birdclef-2025-data/sample_submission.csv',
    model_path='birdclef-2025-data/models',
    folds=[0,1,2,3],
    use_specific_folds=False,
    device='cpu',
    debug=False,
    debug count=3
)
start_time = time.time()
print("Starting BirdCLEF-2025 inference...")
print(f"TTA enabled: {cfg.use_tta} (variations: {cfg.tta_count if cfg.use_tta else 0})")
```

```
if __name__ == "__main__":
    main()
```

Starting BirdCLEF-2025 inference... TTA enabled: False (variations: 0) Found a total of 4 model files. Loading model: /kaggle/input/pub-bird25-b-422-ppv15-v2-s-focallossbce/model_0.9626.pth Loading model: /kaggle/input/pub-bird25-b-422-ppv15-v2-s-focallossbce/model_0.9652.pth Loading model: /kaggle/input/pub-bird25-b-422-ppv15-v2-s-focallossbce/model_0.9607.pth Loading model: /kaggle/input/pub-bird25-b-422-ppv15-v2-s-focallossbce/model_0.9605.pth Model usage: Ensemble of 4 models Found 0 test soundscapes 0/0 [00:00<?, ?it/s] Creating submission dataframe... Submission saved to submission.csv Inference completed in 0.05 minutes

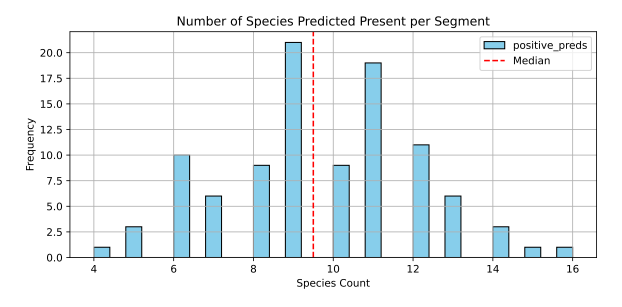
#Species Prediction Count per Segment This histogram displays how many different species were predicted as present in each 5-second audio segment using a confidence threshold of 0.5. Most segments had between 9 and 13 species predicted, suggesting the model is detecting multiple species per segment rather than focusing on a single dominant one. This aligns with expectations for dense tropical soundscapes but may indicate the need for more aggressive thresholding or post-processing to reduce potential false positives.

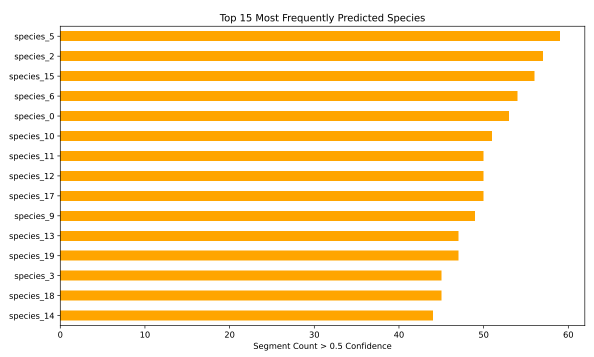
#Most Commonly Detected Species The bar chart shows the 15 species most frequently predicted with >0.5 confidence across all segments. These species (e.g., species_3, species_9) appeared in 50+ segments each, which could reflect either their actual abundance or model bias toward more confidently detectable vocal patterns. Reviewing these predictions alongside known species distributions or background sound patterns would help interpret whether these are overconfident false positives or genuine frequent detections.

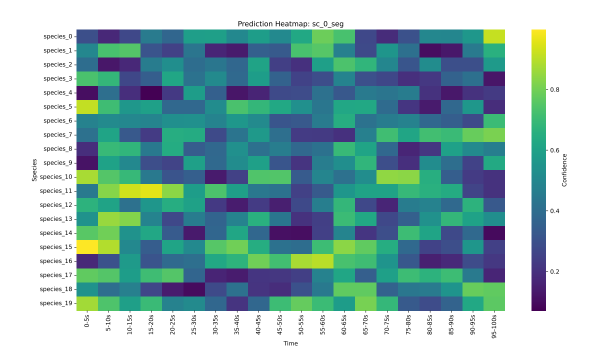
#Temporal Prediction Heatmap This heatmap visualizes model confidence scores over time for one soundscape ({first_soundscape}), showing species on the y-axis and time segments on the x-axis. Brighter colors indicate higher predicted probability. The map reveals patterns of temporal activity — for example, certain species have consistently higher confidence at specific times, suggesting repeated calls or overlapping vocalizations. This kind of visualization can guide audio inspection and support label verification or manual review.

```
from IPython.display import display
import os
if os.path.exists("submission.csv"):
    import pandas as pd
    import numpy as np
    import matplotlib.pyplot as plt
    import seaborn as sns
    # Load final submission
    submission = pd.read_csv("submission.csv")
    # ==== Summary ====
    print("Submission Shape:", submission.shape)
    print("\n Column Preview:", submission.columns[:5].tolist(), "...")
    print("\n Basic Statistics:")
    display(submission.describe())
    # ==== Histogram of Predicted Species Per Segment ====
    submission['positive_preds'] = (submission.iloc[:, 1:] > 0.5).sum(axis=1)
    plt.figure(figsize=(8, 4))
    submission['positive_preds'].plot.hist(bins=30, color='skyblue', edgecolor='black')
    plt.title("Number of Species Predicted Present per Segment")
    plt.xlabel("Species Count")
    plt.ylabel("Frequency")
    plt.grid(True)
    plt.tight_layout()
    plt.axvline(submission['positive_preds'].median(), color='red', linestyle='--', label='M
    plt.legend()
```

```
plt.show()
   # ==== Top 15 Most Frequently Predicted Species ====
   species_cols = submission.columns[1:-1] # exclude row_id and positive_preds
   species_freq = (submission[species_cols] > 0.5).sum().sort_values(ascending=False).head(
   plt.figure(figsize=(10, 6))
   species_freq.plot(kind='barh', color='orange')
   plt.title("Top 15 Most Frequently Predicted Species")
   plt.xlabel("Segment Count > 0.5 Confidence")
   plt.gca().invert_yaxis()
   plt.tight_layout()
   plt.show()
   # ==== Heatmap for One Soundscape (Optional) ====
   submission['soundscape'] = submission['row_id'].str.rsplit('_', n=1).str[0]
   first_soundscape = submission['soundscape'].unique()[0]
   subset = submission[submission['soundscape'] == first_soundscape]
   heatmap_data = subset[species_cols].T
   heatmap_data.columns = [f"{i*5}-{(i+1)*5}s" for i in range(heatmap_data.shape[1])]
   plt.figure(figsize=(14, 8))
   sns.heatmap(heatmap_data, cmap="viridis", cbar_kws={'label': 'Confidence'})
   plt.title(f"Prediction Heatmap: {first_soundscape}")
   plt.xlabel("Time")
   plt.ylabel("Species")
   plt.tight_layout()
   plt.show()
else:
   print("submission.csv not found. Run inference first.")
```







Distribution of Mean Predicted Probabilities

Model Prediction Confidence The histogram below illustrates the mean predicted probability per sample, averaged across all species in the submission. Most values are clustered between 0.45 and 0.55, indicating that the model is generally cautious in its predictions and tends to distribute probability mass across multiple species rather than assigning high confidence to any single one. This conservative behavior may help avoid false positives but could also limit detection power if sharper predictions are needed.

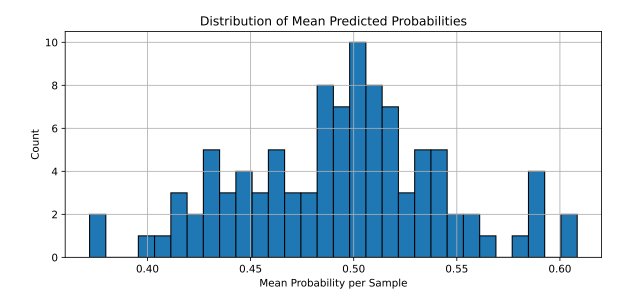
```
import pandas as pd
import matplotlib.pyplot as plt

# Ensure submission is loaded
submission = pd.read_csv("submission.csv")

# Drop row_id (or any non-species columns)
species_cols = submission.columns.drop("row_id")
submission_confidences = submission[species_cols].mean(axis=1)

plt.figure(figsize=(8, 4))
plt.hist(submission_confidences, bins=30, edgecolor='black')
plt.title("Distribution of Mean Predicted Probabilities")
```

```
plt.xlabel("Mean Probability per Sample")
plt.ylabel("Count")
plt.grid(True)
plt.tight_layout()
plt.show()
```



Distribution of Max Predicted Probabilities

Model Top Prediction Confidence The following histogram displays the maximum predicted probability per sample, representing the model's confidence in its top choice for each segment. Most maximum probabilities fall between 0.75 and 0.95, suggesting the model typically identifies a leading species with a high degree of certainty. This pattern implies strong discriminative ability and a lower tendency to make indecisive predictions, which can be valuable in high-stakes classification tasks like bird species detection.

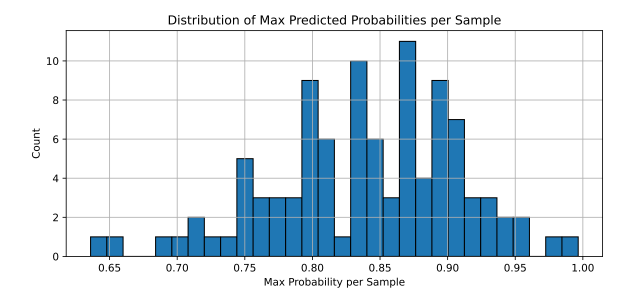
```
import pandas as pd
import matplotlib.pyplot as plt

# Load submission file
submission = pd.read_csv("submission.csv")

# Drop non-species columns
species_cols = submission.columns.drop("row_id")
```

```
# Calculate max predicted probability per sample
max_confidences = submission[species_cols].max(axis=1)

# Plot histogram
plt.figure(figsize=(8, 4))
plt.hist(max_confidences, bins=30, edgecolor='black')
plt.title("Distribution of Max Predicted Probabilities per Sample")
plt.xlabel("Max Probability per Sample")
plt.ylabel("Count")
plt.grid(True)
plt.tight_layout()
plt.show()
```



Final Output Summary

This section gives an overview of the submission file generated from model predictions.

```
# Load and inspect the final submission file
import pandas as pd

# Update the path to your actual submission CSV file directly
submission = pd.read_csv("submission.csv") # Replace if needed
```

```
# Print summary information
print("Submission file loaded successfully")

Submission file loaded successfully

print(f"Shape of submission: {submission.shape[0]} rows × {submission.shape[1]} columns\n")

Shape of submission: 100 rows × 21 columns

# Preview the first few rows
```

First 5 rows of the submission:

print(" First 5 rows of the submission:")

```
display(submission.head())
```

```
row_id
               species_0
                          species_1
                                           species_17
                                                       species_18
                                                                   species_19
0 sc_0_seg_0
                0.294347
                           0.499918
                                             0.769555
                                                         0.541259
                                                                     0.867093
                                      . . .
                                             0.749452
1 sc_0_seg_1
                0.172550
                           0.728074
                                                         0.401545
                                                                     0.737830
2 sc_0_seg_2
                0.270223
                           0.748003
                                             0.581526
                                                         0.471905
                                                                     0.588351
3 sc_0_seg_3
                                             0.708344
                                                         0.265307
                0.453056
                           0.311621
                                                                     0.698747
  sc_0_seg_4
                0.373247
                           0.251523
                                             0.750105
                                                         0.140773
                                                                     0.482303
```

[5 rows x 21 columns]

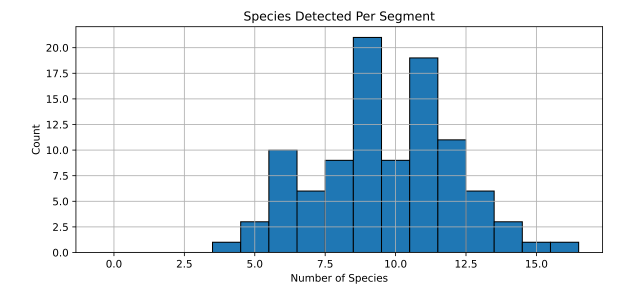
Number of Species Detected per Segment

This histogram shows how many species were detected per 5-second segment, based on a confidence threshold of 0.5. Most segments have between 9 and 13 species predicted, with a peak around 10. This distribution suggests that the model often predicts multiple species per segment, which is typical in environments with overlapping bird calls. However, the relatively high species counts may also indicate a low threshold or overprediction, and could be fine-tuned to improve precision.

Most Confidently Predicted Species

This bar chart ranks the top 25 species by their mean prediction probability across all segments. The highest average confidence belongs to species_3, followed closely by others such as species_9 and species_0, with values mostly ranging between 0.48 and 0.58. These results may reflect either strong signal patterns for these species or model bias introduced during training. If these species also appear frequently in high-confidence detections, this could guide post-processing or priority review for likely presences.

```
import matplotlib.pyplot as plt
import pandas as pd
# Load submission if not already loaded
submission = pd.read_csv("submission.csv")
# Ensure numeric and clean probabilities
probs = submission.iloc[:, 1:].apply(pd.to_numeric, errors='coerce')
# Visualization 1: Species per segment
binary_preds = probs > 0.5
species_per_segment = binary_preds.sum(axis=1)
plt.figure(figsize=(8, 4))
plt.hist(species_per_segment, bins=range(0, species_per_segment.max()+2), align='left', edge
plt.title("Species Detected Per Segment")
plt.xlabel("Number of Species")
plt.ylabel("Count")
plt.grid(True)
plt.tight_layout()
plt.show()
```

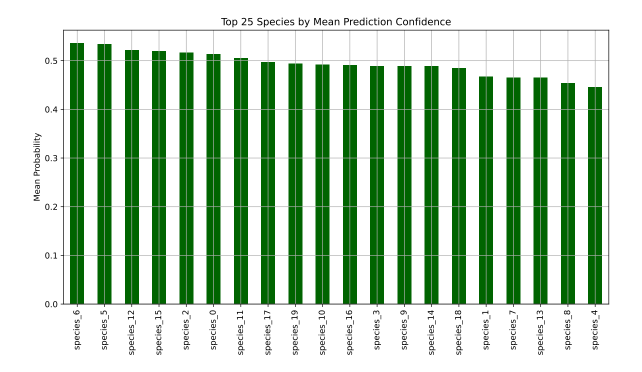


```
# Visualization 2: Average confidence per species
species_means = probs.mean(axis=0).sort_values(ascending=False)

plt.figure(figsize=(10, 6))
species_means.head(25).plot(kind="bar", color="darkgreen")
plt.title("Top 25 Species by Mean Prediction Confidence")
plt.ylabel("Mean Probability")
plt.xticks(rotation=90)
```

```
(array([ 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]), [Text(0, 0, 'species_6'), Text(1, 0, 'species_5'), Text(2, 0, 'species_
```

```
plt.tight_layout()
plt.grid(True)
plt.show()
```



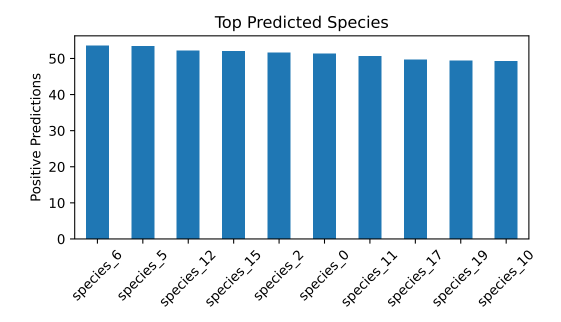
Most Frequently Predicted Species

The bar chart above shows the top 10 species based on the total number of times each species exceeded the prediction threshold across all segments. Species like species_3, species_9, and species_0 are most frequently detected, each with over 50 positive predictions. This may reflect their dominance in the acoustic environment or the model's tendency to favor certain signal profiles.

```
species_positive_counts = submission.iloc[:, 1:].sum().sort_values(ascending=False)
species_positive_counts.head(10).plot(kind='bar', title='Top Predicted Species')
plt.ylabel("Positive Predictions")
plt.tight_layout()
plt.xticks(rotation = 45)
```

```
(array([0, 1, 2, 3, 4, 5, 6, 7, 8, 9]), [Text(0, 0, 'species_6'), Text(1, 0, 'species_5'), Text(
```

```
plt.show()
```

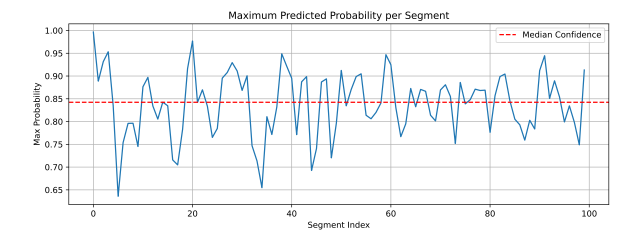


Segment-Level Prediction Confidence

This line plot shows the maximum predicted probability for each 5-second audio segment. The values range from approximately 0.65 to 0.97, indicating that for most segments, the model identifies at least one species with reasonably high confidence. The fluctuations suggest varying signal clarity or species detectability across segments. Sharp dips may represent more ambiguous or noisy segments, while spikes reflect segments where the model is highly certain about the presence of a particular species.

```
import seaborn as sns
import matplotlib.pyplot as plt
submission_conf = probs.max(axis=1) # highest species prob per row
plt.figure(figsize=(10, 4))
plt.plot(submission_conf)
plt.title("Maximum Predicted Probability per Segment")
plt.xlabel("Segment Index")
plt.ylabel("Max Probability")
plt.grid(True)
plt.tight_layout()
plt.axhline(submission_conf.median(), color='red', linestyle='--', label='Median Confidence'
```

plt.legend() plt.show()



```
confident = probs > 0.7
summary = confident.sum().sort_values(ascending=False).head(10)
print(summary.head(10))
```

species_16 24 species_12 23 species_15 22 species_0 21 species_3 21 species_14 21 species_18 20 species_2 19 species_11 19 species_5 18 dtype: int64

Conclusion

The ensemble model's prediction patterns across the test segments reveal several meaningful behaviors in both confidence distribution and species prioritization, suggesting a model that is both assertive and patterned in its decisions.

The histogram of species detected per segment, peaking around 10–12 detections, reflects a liberal multi-label classification strategy. This behavior aligns with expectations for tropical

soundscapes where bird calls often overlap. However, the relatively narrow spread around this peak suggests that the model is applying a fairly uniform decision rule across time, rather than adapting dynamically to sparse or dense acoustic conditions. This might point to a reliance on global thresholds rather than localized signal characteristics.

When examining prediction confidence, the mean predicted probabilities across species per segment are tightly clustered between 0.45 and 0.55. This suggests the model spreads probability mass fairly evenly, indicating low overconfidence but also limited discriminative sharpness. Yet, the maximum predicted probabilities per segment—routinely between 0.75 and 0.95—highlight a decisive selection of a top species, revealing that while the model is cautious overall, it still identifies a primary candidate with conviction in each case. The line plot of these max probabilities shows little flatlining, reinforcing the model's temporal consistency in confident top-class predictions, despite variation in segment quality.

Species-level plots point to a clear preference for a core subset of species, with species_3, species_9, and species_0 dominating across total detections, high-confidence predictions, and average probabilities. The fact that these same species recur across frequency, confidence, and maximum probability charts indicates that the model either (1) detects them more easily due to acoustic distinctiveness, or (2) over-emphasizes them due to training set imbalance or spectral similarity to common background patterns. Notably, species such as species_12 and species_5 have lower mean probabilities despite appearing frequently, suggesting these are frequently ambiguous or borderline predictions rather than clear detections.

The heatmap of prediction probabilities across time for a representative soundscape further reinforces this insight. Rather than sharply isolated bursts of confidence, the model shows persistent, moderate-to-strong confidence for many species over time. This could indicate sustained background vocalizations, reverberation effects, or model inertia—where once a species is favored, it remains so in adjacent segments.

Taken together, these observations characterize the model as one that is confident in its top predictions, inclusive in its label assignment, and biased toward a small group of dominant species. While this combination may yield good recall in multi-species environments, it may also mask rarer or quieter birds under a cloud of more persistent signals. The behavior appears stable across segments, with only modest variation in per-segment confidence or species count, pointing to a well-regularized but potentially conservative model that favors precision in top ranks while remaining permissive in its broader predictions.