BiSSe - Binary State Speciation and Extinction

The model described is a Birth-Death model with two interacting species and the possibility of transitions between them. This model captures the dynamics of two species populations over time, incorporating birth, death, and transition rates.

Parameters

- λ_1, λ_2 : Birth rates of species 1 and species 2, respectively.
- μ_1, μ_2 : Death rates of species 1 and species 2, respectively.
- p_{12}, p_{21} : Transition rates from species 1 to species 2 and from species 2 to species 1, respectively.
- ini_1, ini_2 : Initial populations of species 1 and species 2, respectively.
- T: Maximum time for the simulation.

Model Dynamics

- 1. **Initialization**: The initial populations of species 1 and species 2 are set based on the given parameters ini1 and ini2. The current time is initialized to zero.
- 2. **Event Simulation**: The process continues in a loop until the current time exceeds the maximum time T or both species' populations become zero.
 - Total Rate Calculation: At each step, the total rate of events is calculated as the sum of all possible events' rates:

$$ext{total_rate} = n_1(\lambda_1 + \mu_1 + p_{12}) + n_2(\lambda_2 + \mu_2 + p_{21})$$

where n_1 and n_2 are the current populations of species 1 and species 2, respectively.

• Event Time Sampling: The time until the next event is sampled from an exponential distribution with the rate parameter total_rate.

• **Event Type Sampling**: The type of event is determined by sampling from a discrete distribution with probabilities proportional to the rates of each event:

$$\text{event} \setminus \text{probs} = \left[\frac{n_1 \lambda_1}{\text{total} \setminus \text{rate}}, \frac{n_2 \lambda_2}{\text{total} \setminus \text{rate}}, \frac{n_1 \mu_1}{\text{total} \setminus \text{rate}}, \frac{n_2 \mu_2}{\text{total} \setminus \text{rate}}, \frac{n_1 p_{12}}{\text{total} \setminus \text{rate}}, \frac{n_2 p_{21}}{\text{total} \setminus \text{rate}} \right]$$

- Event Execution: Based on the sampled event type, the populations are updated accordingly:
 - Birth of species 1: $(n_1 \leftarrow n_1 + 1)$
 - Birth of species 2: $(n_2 \leftarrow n_2 + 1)$
 - Death of species 1: $(n_1 \leftarrow n_1 1)$
 - Death of species 2: $(n_2 \leftarrow n_2 1)$
 - lacktriangle Transition from species 1 to species 2: $(n_1 \leftarrow n_1 1), (n_2 \leftarrow n_2 + 1)$
 - lacktriangle Transition from species 2 to species 1: $(n_2 \leftarrow n_2 1), (n_1 \leftarrow n_1 + 1)$
- Event Recording: Each event, along with the current time and updated populations, is recorded.
- 3. **Termination**: The process stops when the current time exceeds the maximum time T or both species' populations reach zero.

Output

The function returns a list of events, each represented as a tuple (time, n1, n2), where time is the time of the event, and n1 and n2 are the populations of species 1 and species 2 after the event.

$$ext{total} \setminus ext{rate} = n_1(\lambda_1 + \mu_1 + p_{12}) + n_2(\lambda_2 + \mu_2 + p_{21})$$

```
In []: import numpy as np

def bisse(lam1, lam2, mu1, mu2, p12, p21, ini1, ini2, T, limit_event_size = 1000):
```

```
n1 = ini1.copy()
n2 = ini2.copy()
current time = 0
events = []
events_list = np.array([1,2,3,4,5,6])
final T = T
while current time < T:</pre>
    total population = n1 + n2
    if total population == 0:
        break
    if len(events) > limit event size:
        final_T = current_time
        break
    total rate = n1*(lam1+mu1+p12) + n2*(lam2+mu2+p21)
    sampled_time = np.random.exponential(1/total_rate)
    current time += sampled time
    if current_time > T:
        break
    event probs = n_array([n1*lam1, n2*lam2, n1*mu1, n2*mu2, n1*p12, n2*p21])/total rate
    event = np.random.choice(events list, p=event probs)
    match event:
        case 1: # specie 1 gives birth
            n1 += 1
        case 2: # specie 2 gives birth
            n2 += 1
        case 3: # specie 1 dies
            n1 -= 1
        case 4: # specie 2 dies
            n2 -= 1
```

```
case 5: # specie 1 transitions to specie 2
    n1 -= 1
    n2 += 1

case 6: # specie 2 transitions to specie 1
    n2 -= 1
    n1 += 1

case _:
    raise ValueError("Invalid event")

events.append((current_time, n1, n2))

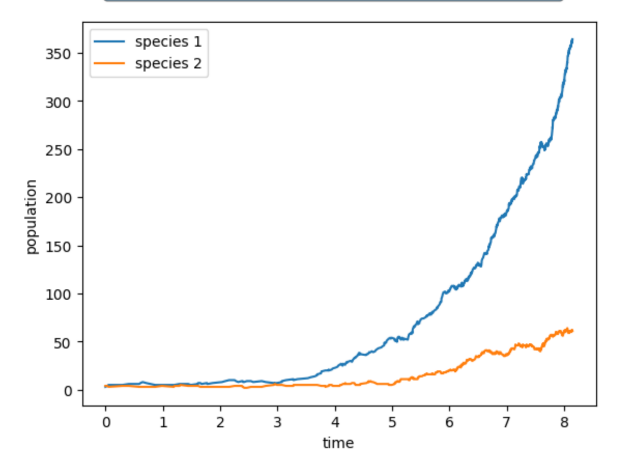
return final_T, events
```

We randomize the paramaters and make an experiment by computing BiSSe and then ploting the evolution of the species over time

```
In [ ]: # parameters
        lam1, lam2 = np.random.uniform(0, 1, size=2)
        mu1, mu2 = np.random.uniform(0, 0.8, size=2)
        p12, p21 = np.random.uniform(0, 0.5, size=2)
        max time = 10
        \max num initial population = 5
        ini1, ini2 = np.random.randint(0, max num initial population, size=2)
        final T, events = bisse(lam1, lam2, mu1, mu2, p12, p21, ini1, ini2, max time)
        nodes = [(0, ini1, ini2)] + events
        # plot each of the species in the same figure
        import matplotlib.pyplot as plt
        fig, ax = plt.subplots()
        ax.plot([node[0] for node in nodes], [node[1] for node in nodes], label='species 1')
        ax.plot([node[0] for node in nodes], [node[2] for node in nodes], label='species 2')
        ax.set xlabel('time')
        ax.set_ylabel('population')
        rates_info = f'lam1={lam1:.2f}, lam2={lam2:.2f}, mu1={mu1:.2f}, mu2={mu2:.2f}, p12={p12:.2f}, p21={p21:.2f}'
```

ax.text(0.05, 1.1, rates_info, transform=ax.transAxes, fontsize=9, verticalalignment='top', bbox=dict(boxstyle="
ax.legend()
plt.show()

lam1=0.93, lam2=0.66, mu1=0.28, mu2=0.61, p12=0.17, p21=0.34



Function Description

The generate_data function simulates the dynamics of two interacting species over multiple iterations and collects the resulting

data. I his function leverages the Birth-Death model with transition rates between the species to generate the data points.

Parameters

- num_data_points: The number of data points to generate.
- max_lamb_rate: The maximum value for the birth rates λ_1 and λ_2 .
- max_mniu_rate: The maximum value for the death rates μ_1 and μ_2 .
- max_num_initial_population: The maximum initial population size for both species.
- max_time: The maximum simulation time for each iteration.

Function Dynamics

- 1. **Initialization**: Two empty lists, X and Y, are created to store the input parameters and the resulting populations, respectively.
- 2. Loop through Data Points: For each data point:
 - Randomly sample birth rates λ_1 and λ_2 from a uniform distribution between 0 and max_lamb_rate.
 - Randomly sample death rates μ_1 and μ_2 from a uniform distribution between 0 and max_mniu_rate.
 - Randomly sample transition rates p_{12} and p_{21} from a uniform distribution between 0 and 1.
 - Randomly sample a simulation time time from a uniform distribution between 0 and max_time.
 - Randomly sample initial populations ini1 and ini2 from an integer uniform distribution between 0 and max_num_initial_population.
- 3. **Simulation**: For each set of sampled parameters, the bisse function is called to simulate the population dynamics over the sampled time period.
- 4. **Event Recording**: The populations of species 1 and species 2 at the end of the simulation are recorded. If no events occurred during the simulation, the initial populations are used.
- 5. Data Collection: The sampled parameters and the resulting populations are appended to the lists X and Y.
- 6 Return Values: The function returns two NumPv arrays X and Y where X contains the input parameters for each data point

and Y contains the resulting populations of species 1 and species 2.

```
In []: from tqdm.notebook import tqdm

def generate_data(num_data_points, max_lamb_rate, max_mniu_rate, max_num_initial_population, max_time):
    X = []
    Y = []

    for _ in tqdm(range(num_data_points)):
        lam1, lam2 = np.random.uniform(0, max_lamb_rate, size=2)
        mu1, mu2 = np.random.uniform(0, max_mniu_rate, size=2)
        p12, p21 = np.random.uniform(0, 1, size=2)
        time = np.random.uniform(0, max_time)
        ini1, ini2 = np.random.randint(0, max_num_initial_population, size=2)

    final_T, events = bisse(lam1, lam2, mu1, mu2, p12, p21, ini1, ini2, time)
    _, num_specie1, num_specie2 = (0, ini1, ini2) if len(events) == 0 else events[-1]
    X.append([lam1, lam2, mu1, mu2, p12, p21, ini1, ini2, final_T])
    Y.append([num_specie1, num_specie2])

return np.array(X), np.array(Y)
```

Data Generation and Splitting

Data Generation

The data generation process involves simulating the dynamics of two interacting species over a large number of iterations using the generate_data function. The parameters for this process are as follows:

• num_data_points: (64 imes 1250)

- max_lamb_rate: 1
- max mniu rate: 1
- max_num_initial_population: 5
- max_time: 10

The function generate_data is called with these parameters to produce the input data X and the corresponding output data Y.

Note:

• We consider a small time frame of max time 10 because if sampled lambdas are substantially greater than the sampled mnius, then the growth of the population is exponential and this will become computationally unfeasible.

```
In []: import pandas as pd
        num_data_points = 64*1250
        max lamb rate = 1
        max mniu rate = 1
        \max num initial population = 5
        max time = 10
        X, Y = generate data(num data points, max lamb rate, max mniu rate, max num initial population, max time)
        df_X = pd.DataFrame(X, columns=['lam1', 'lam2', 'mu1', 'mu2', 'p12', 'p21', 'ini1', 'ini2', 'final_T'])
        df Y = pd.DataFrame(Y, columns=['num specie1', 'num specie2'])
        df = pd.concat([df_X, df_Y], axis=1)
        from sklearn.model selection import train test split
        train set, test set = train test split(df, test size=0.1)
        train set, val set = train test split(train set, test size=0.2)
                      | 0/80000 [00:00<?, ?it/s]
         0%|
In [ ]: train set.corr()
```

]:		lam1	lam2	mu1	mu2	p12	p21	ini1	ini2	final_T	num_speci
	lam1	1.000000	-0.000070	0.000534	0.001622	0.003461	0.005749	0.004263	-0.004258	-0.005277	0.20586
	lam2	-0.000070	1.000000	-0.010784	-0.000992	0.000831	-0.004449	-0.004942	-0.015801	-0.010984	0.0740
	mu1	0.000534	-0.010784	1.000000	0.009937	0.007361	0.000910	0.000710	-0.005827	0.006041	-0.2206
	mu2	0.001622	-0.000992	0.009937	1.000000	0.008823	-0.004641	0.005835	-0.007039	0.002614	-0.0868
	p12	0.003461	0.000831	0.007361	0.008823	1.000000	0.000222	0.007700	-0.002718	-0.005402	-0.1243
	p21	0.005749	-0.004449	0.000910	-0.004641	0.000222	1.000000	0.005385	-0.001673	-0.000859	0.05620
	ini1	0.004263	-0.004942	0.000710	0.005835	0.007700	0.005385	1.000000	0.002921	-0.010974	0.07746
	ini2	-0.004258	-0.015801	-0.005827	-0.007039	-0.002718	-0.001673	0.002921	1.000000	-0.005296	0.05969
	final_T	-0.005277	-0.010984	0.006041	0.002614	-0.005402	-0.000859	-0.010974	-0.005296	1.000000	0.1577
	num_specie1	0.205866	0.074091	-0.220685	-0.086814	-0.124361	0.056205	0.077465	0.059690	0.157798	1.00000
	num_specie2	0.069280	0.208831	-0.087431	-0.222439	0.047301	-0.123966	0.051084	0.086869	0.153907	0.31869

```
In []: train_X = train_set.drop(columns=['num_specie1', 'num_specie2'])
    train_Y = train_set[['num_specie1', 'num_specie2']].copy()

val_X = val_set.drop(columns=['num_specie1', 'num_specie2'])
    val_Y = val_set[['num_specie1', 'num_specie2']].copy()

test_X = test_set.drop(columns=['num_specie1', 'num_specie2'])
    test_Y = test_set[['num_specie1', 'num_specie2']].copy()

#see missing values
    print(train_X.isnull().sum())
    print(val_X.isnull().sum())

print(val_X.isnull().sum())

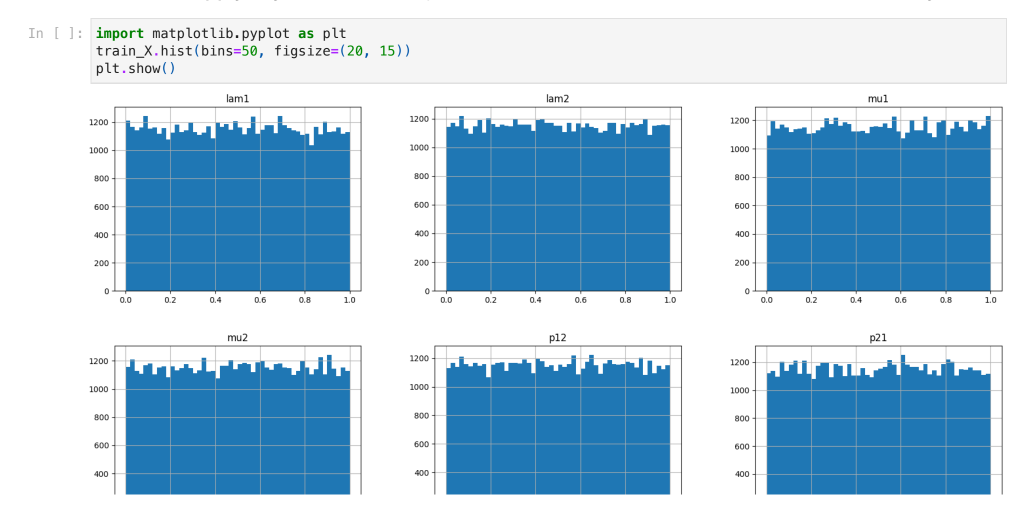
print(val_Y.isnull().sum())
```

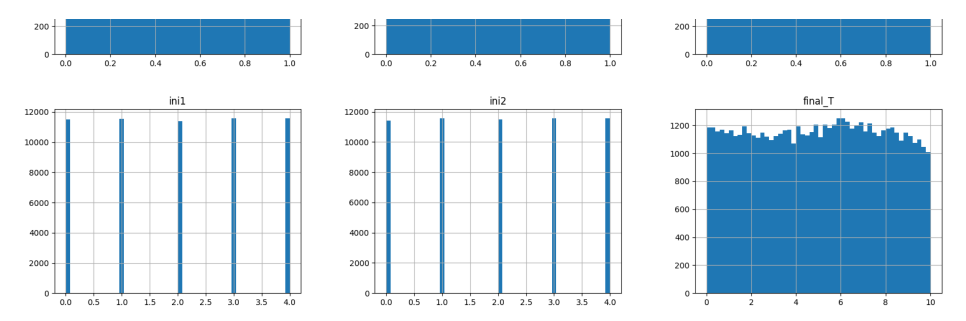
Out[]

```
pi iiic (vac_i isiiucc() isum())
 print(test_X.isnull().sum())
 print(test_Y.isnull().sum())
lam1
           0
lam2
           0
mu1
           0
mu2
p12
p21
ini1
ini2
final_T
dtype: int64
num_specie1
               0
               0
num_specie2
dtype: int64
lam1
           0
lam2
           0
mu1
mu2
p12
p21
ini1
ini2
           0
final_T
dtype: int64
num_specie1
               0
num_specie2
dtype: int64
lam1
lam2
           0
mu1
mu2
p12
p21
           0
ini1
           a
```

ini2 0
final_T 0
dtype: int64
num_specie1
num_specie2
dtype: int64

Before we apply any transformation, we should see if distribution of each attribute is heavy tail.





From above we can see the distribution is uniform

Now we can start building the pipeline

```
Out[]: array([[ 0.85966154, 0.86625492, -0.88649332, ..., -1.
                     , 0.79953382],
                -0.5
               [ 0.89198854, 0.18058109, 0.04330034, ..., 0.5
                     , -0.6881473 ],
               [-0.2677207, 0.66436848, -0.34219325, ..., 1.
                          . -0.96937705],
                -0.5
               [-0.94178962, -0.08779862, 0.18552385, ..., -0.5
                    , -0.09070316],
               [-0.67850873, -0.86402238, -0.60835774, ..., 0.5]
                          , -0.73872887],
                 0.5
               [0.10871981, -0.81089319, -0.97158014, ..., -0.5]
                 0.5
                           , -0.60328665]])
In []: val X prepared = pipeline.transform(val X)
        test X prepared = pipeline.transform(test X)
        train Y prepared = train Y.to numpy()
        val Y prepared = val Y.to numpy()
        test Y prepared = test Y.to numpy()
```

Neural Network Model Training and Visualization Neural Network Model Architecture

The code defines a neural network model using TensorFlow's Keras API. The model architecture consists of:

- Input Layer: Defined by the shape of X_train[0], which corresponds to the shape of the input data.
- **Dense Layers**: Four hidden layers with 16, 26, 18, and 8 neurons respectively, each using ReLU (Rectified Linear Unit) activation function.
- Output Layer: An output layer with neurons equal to the number of outputs (Y_train[0].shape[0]), which predicts the populations of species 1 and species 2.

Callback

• **Early Stopping**: A callback (ea_callback) is used to monitor the validation loss (val_loss). Training will stop early if the validation loss does not improve for 5 consecutive epochs (patience=5). The model will restore the weights that give the best validation loss (restore_best_weights=True).

Model Compilation

The model is compiled using the Adam optimizer (optimizer='adam') and mean squared error (loss='mse') as the loss function. The accuracy metric is used for evaluation (metrics=['accuracy']).

Model Training

The model.fit method is called to train the model:

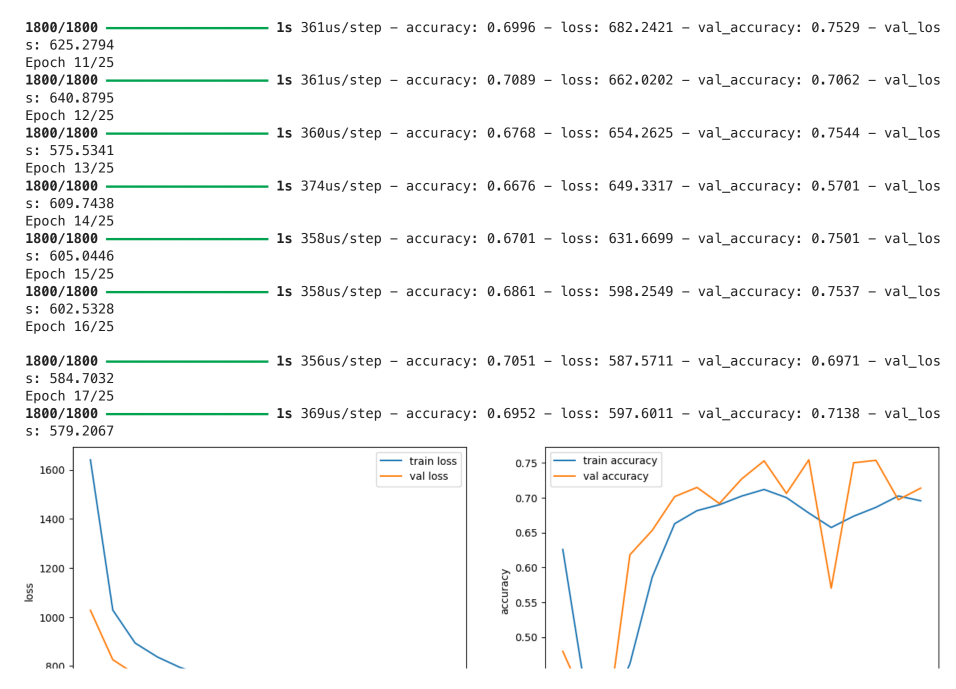
- X_train, Y_train: Training data and labels.
- epochs: Number of epochs set to 25.
- batch_size: Batch size set to 32.
- validation_data: Validation data and labels provided as (X_val, Y_val).
- callbacks: Early stopping callback (ea_callback) is passed to monitor validation loss during training.

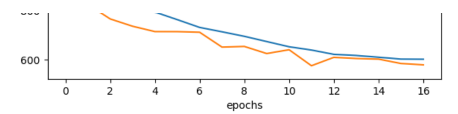
Training History Visualization

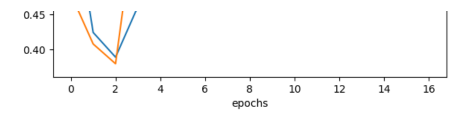
After training, the accuracy and validation accuracy over epochs are plotted using Matplotlib to visualize the model's performance.

```
-0.5
                            , -0.9693//05],
                ...,
                [-0.94178962, -0.08779862, 0.18552385, ..., -0.5]
                            , -0.09070316],
                 0.
                [-0.67850873, -0.86402238, -0.60835774, \ldots, 0.5]
                 0.5
                            , -0.73872887],
                [ 0.10871981, -0.81089319, -0.97158014, ..., -0.5 ]
                            , -0.60328665]])
                 0.5
In [ ]: train_Y_prepared[0].shape[0]
Out[]: 2
In [ ]: import tensorflow as tf
        model = tf.keras.Sequential([
            tf.keras.layers.InputLayer(train X prepared[0].shape),
            tf.keras.layers.Dense(16, activation='relu'),
            tf.keras.layers.Dense(26, activation='relu'),
            tf.keras.layers.Dense(18, activation='relu'),
            tf.keras.layers.Dense(8, activation='relu'),
            tf.keras.layers.Dense(train Y prepared[0].shape[0])
        1)
        ea callback = tf.keras.callbacks.EarlyStopping(monitor='val loss', patience=5, restore best weights=True)
        model.compile(optimizer='adam', loss='mse', metrics=['accuracy'])
        history = model.fit(train X prepared, train Y, epochs=25, batch size=32, validation data=(val X prepared, val Y)
        fig, ax = plt.subplots(1, 2, figsize=(15, 5))
        ax[0].plot(history.history['loss'], label='train loss')
        ax[0].plot(history.history['val loss'], label='val loss')
        ax[0].set xlabel('epochs')
        ax[0].set ylabel('loss')
        ax[0].legend()
```

```
ax[1].plot(history.history['accuracy'], label='train accuracy')
 ax[1].plot(history.history['val accuracy'], label='val accuracy')
 ax[1].set xlabel('epochs')
 ax[1].set ylabel('accuracy')
 ax[1].legend()
 plt.show()
Epoch 1/25
                            — 1s 404us/step - accuracy: 0.6331 - loss: 2018.6576 - val_accuracy: 0.4796 - val_lo
1800/1800 —
ss: 1027.3483
Epoch 2/25
                          ——— 1s 360us/step - accuracy: 0.4450 - loss: 1072.3293 - val_accuracy: 0.4078 - val_lo
1800/1800 -
ss: 826.0690
Epoch 3/25
                          ——— 1s 369us/step – accuracy: 0.3934 – loss: 918.1927 – val_accuracy: 0.3794 – val_los
1800/1800 -
s: 766.6844
Epoch 4/25
                            — 1s 357us/step - accuracy: 0.4228 - loss: 825.2649 - val accuracy: 0.6181 - val los
1800/1800 -
s: 736.8059
Epoch 5/25
                            — 1s 355us/step - accuracy: 0.5710 - loss: 749.8409 - val accuracy: 0.6532 - val los
1800/1800 -
s: 714.8970
Epoch 6/25
                            — 1s 357us/step - accuracy: 0.6494 - loss: 768.8055 - val accuracy: 0.7015 - val los
1800/1800 -
s: 714.7110
Epoch 7/25
1800/1800 -
                            — 1s 358us/step - accuracy: 0.6748 - loss: 726.2449 - val accuracy: 0.7148 - val los
s: 712,2497
Epoch 8/25
                             — 1s 372us/step - accuracy: 0.6778 - loss: 713.0627 - val_accuracy: 0.6917 - val los
1800/1800 -
s: 651.6809
Epoch 9/25
                            — 1s 365us/step - accuracy: 0.7146 - loss: 690.7686 - val_accuracy: 0.7270 - val_los
1800/1800 -
s: 654,3060
Epoch 10/25
```







Model Evaluation on Test Data

Model Evaluation

To evaluate the trained neural network model on the test data (X_test and Y_test), the model.evaluate method is used. This method computes the loss and metrics (accuracy in this case) on the test set.

Saving the model

We can observe that our current model has an accuracy of 72.2% on the test set. In other words, given a vector space of parameters within the bounds previously defined, we can predict the number of of each species with an accuracy of 72.2%. We can proceed and save the model for future use.

```
In []: # save the model
model.save('bisse_model.keras')
```