Deep Multi-Species Embedding

Di Chen1 Yexiang Xue2 Daniel Fink3 Shuo Chen4 Carla P. Gomes5
1 2 4 5Department of Computer Science, Cornell University, Ithaca, NY , USA
3Cornell Lab of Ornithology, Ithaca, NY , USA
{1dc874,3df36}@cornell.edu {2yexiang,4shuochen,5gomes}@cs.cornell.edu

Abstract

Understanding how species are distributed across landscapes over time is a fundamental question in biodiversity research. Unfortunately, most species distribution models only target a single species at a time, despite strong ecological evidence that species are not independently distributed. We propose Deep Multi-Species Embedding (DMSE), which jointly embeds vectors corresponding to multiple species as well as vectors representing environmental covariates into a common high-dimensional feature space via a deep neural network. Applied to bird observational data from the citizen science project eBird, we demonstrate how the DMSE model discovers inter-species relationships to outperform single-species distribution models (random forests and SVMs) as well as competing multi-label models. Additionally, we demonstrate the benefit of using a deep neural network to extract features within the embedding and show how they improve the predictive performance of species distribution modelling. An important domain contribution of the DMSE model is the ability to discover and describe species interactions while simultaneously learning the shared habitat preferences among species. As an additional contribution, we provide a graphical embedding of hundreds of bird species in the Northeast US.

1 Introduction

Understanding the spatial distribution of species and how species interact with each other and their environment is essential for developing science-based conservation plans and ecological research. However, most species distribution models only target a single species at a time [Phillips et al., 2004; Elith and Leathwick, 2009; Fink et al., 2013]. These single-species models ignore the role of species interactions like competition for shared resources (food, territory, etc.). For example, American Robin and Blue Jay are likely to be seen in the same place since the Blue Jay preys on Robin’s eggs or fledglings and sometimes even steals its nest. Therefore, a model that predicts the occupancy of a collection of species instead of modelling each species individually is needed. The most straightforward formulation of a multi-species model [Tsoumakas and Vlahavas, 2007] directly considers the probability of seeing a collection of species. However, this direct approach suffers from combinatorial intractability due to the large number of possible ways to form the collection. As a result, an efficient method of jointly modelling species distribution for large number of species is still lacking.

We propose a novel method called Deep Multi-Species Embedding which can jointly model the distribution of hundreds of species as well as the correlation among species. DMSE jointly embeds multiple species as well as environmental covariates into a high-dimensional feature vector space via a deep neural network. Each embedded vector carries semantic meaning to the modeled entity, and the inner products between them capture the relationships between entities (such as environmental preference or correlation between species).

Applied to eBird bird observational data [Munson et al., 2012], we demonstrate how the DMSE model discovers interspecies relationships to outperform the predictions of single-species distribution models (random forests and SVMs) as well as competing multi-label models. Moreover, as the number of species goes up, the improvement in predictive performance between DMSE models and baseline models keeps widening. Additionally, we demonstrate the benefit of using a deep neural network for feature extraction and show how the features improve the quality of species distribution modelling.

We also show a visualization of the embedding for hundreds of bird species in the Northeast US. It provides an intuitive picture about species shared environmental preferences and the correlations among species, after accounting for those explained by shared environmental preferences. Through this model, we are also able to quantitatively measure many species-species interaction which could only be qualitatively described by ecologists before.
Figure 2: The left graph depicts the independent joint distribution of two species where the color from light yellow to red (better view in color) represents the probability from low to high. The probability mass in each quadrant represents the probability of each two-species co-occurrence. For example, the first quadrant represents the probability that two species occur together. The right graph is derived from the species observation data $D$.

Figure 2: The left graph depicts the independent joint distribution of two species where the color from light yellow to red (better view in color) represents the probability from low to high. The probability mass in each quadrant represents the probability of each two-species co-occurrence. For example, the first quadrant represents the probability that two species occur together. The right graph is derived from the species observation data $D$.

2 Multi-Species Modelling

Our goal is to estimate the joint distribution of multiple species based on the observational data recording the presence or absence of each species at a site. More formally, given a collection of species $\{\text{species}_1, \ldots, \text{species}_n\}$, we would like to estimate the distribution $\Pr(b_i|l_i)$. Here $b_i \in \{0, 1\}^n$ is an indicator for the species co-occurrence of each observation, $b_{i,j} = 1$ if and only if $\text{species}_j$ was detected at site $i$, and $l_i = (f_1, \ldots, f_m)^T$ is an environmental feature vector that contains the values of $m$ environmental covariates (or features) that describe site $i$. To simplify notation, we also use $l_i$ to denote the observation site $i$.

2.1 From One Species to More

Our DMSE method is based on the latent variable formulation of the probit model [Chib and Greenberg, 1998] which is widely used to model binary outcomes. The probit model also has been used by [Pollock et al., 2014] on species distribution modelling. However, their setup is different from ours and can only handle a handful of species.

For the clarity of presentation, we start by describing how to model the distribution of single species using the probit model. For each $\text{species}_i$, we link the occurrence of $\text{species}_i$ at observation site $i$ with a random variable $r_{i,j}$ where the probability that $\text{species}_j$ was detected at observation site $i$ is equal to the probability that $r_{i,j} > 0$, i.e.

$$\Pr(b_{i,j} = 1 | l_i) = \Pr(r_{i,j} > 0)$$

(1)

Here, $r_{i,j}$ follows a normal distribution $N(\mu_{i,j}, \sigma)$ where $\mu_{i,j}$ is a function of $l_i$ and $\sigma$ is fixed to be 1. According to the definition of normal distribution, a positive $\mu_{i,j}$ implies that the $\text{species}_j$ is more likely to be present than absent at site $l_i$ and a negative $\mu_{i,j}$ implies the opposite. Therefore, we can model the distribution of each species by parameterizing $\mu_{i,j}$.

A general approach to model the joint distribution of multiple species is to simply join the distribution of each species assuming each species is independent. For the ease of presentation, we call this kind of joint distribution “independent joint distribution”.

The left graph in the picture above (Fig.2) depicts the independent joint distribution of two species (American Robin and Blue Jay) corresponding to random variable $r_{i,1}$ and $r_{i,2}$. In the graph, the color from light yellow to red represents the probability from low to high and the probability mass in each quadrant represents the probability of each co-occurrence. For example, the probability mass in the first quadrant shows the probability that American Robin and Blue Jay are present together at the observation site $l_i$. The one-dimensional distributions on the graphs’ upper side and left side are the marginal distributions for American Robin and Blue Jay respectively. The red area in each one-dimensional distribution represents the probability of the presence of each species unconditional of the other.

Since the independent joint distribution can not model the correlation between species which widely exists in the real world, we extend the probit model by applying multivariate normal distribution over the $n$-dimensional random variables $r_i = (r_{i,1}, \ldots, r_{i,n})$ i.e.

$$r_i \sim N_n(\mu_i, \Sigma)$$

(2)

where $\mu_i = (\mu_{i,1}, \ldots, \mu_{i,n})^T$ and $\Sigma$ is the covariance matrix. In this way, each random variable $r_{i,j}$ still follows a normal distribution, but we can capture interspecies correlation by parameterizing the covariance matrix $\Sigma$.

As shown in the right graph of Fig 2, we change the covariance between random variable $r_{i,1}$ and $r_{i,2}$ from 0 to a positive number $\rho$, then the joint distribution changes significantly. For example, the probability mass in the first quadrant becomes larger, which means these two species are more likely to be present together. Although we affect the joint distribution of two species by changing the covariance between $r_{i,1}$ and $r_{i,2}$, the marginal distribution of each random variable does not change. This means the probability of the presence of each species, unconditional of the other, is unaffected by the covariance. This property ensures that our model can maintain the predictive capability derived from learning habitat preferences of each species. Meanwhile, it can outperform the independent version when the species distributions are correlated. In addition, if we restrict the
2.2 Deep Multi-Species Embedding

![Diagram of the DMSE framework]

Figure 3: The intuitive visualization of DMSE framework.

In order to estimate the parameters \( \mu \) and \( \Sigma \), we need to first model the species-environment relationship as well as the correlation between species. To achieve this, we first embed each species \( s_j \) with two vectors \( s_j \in R^{d_1} \) and \( \lambda_j \in R^{d_2} \) representing its environmental preference and interactive behavior respectively. Here \( d_1, d_2 \) are the dimensionality of these two vector spaces which are manually set. In our experiments, we set \( d_1 = d_2 = 100. \) Because embedding methods are able to take advantage of high-dimensional representations, it is advantageous to set this parameters to be high, though the methods are not sensitive to the exact values. We choose to model these two characteristics separately instead of embedding to the same vector, because it is not uncommon for groups of species to share similar environment-distribution relationships, but have very different inter-species associations. Thus, by modeling these characteristics separately, DMSE can capitalize on the shared environment-distribution relationships without biasing the inter-species correlation estimates. Moreover, because the environmental features used in the model describe habitat characteristics at a much coarser spatial resolution than that of the inter-species interactions, this model formulation can be seen as a multi-scale approach that shares information at coarse scales at the habitat level while simultaneously allowing fine-scale variation between species.

When it comes to the environmental features, we apply a deep neural network and a projection matrix to embed the low-dimensional raw environmental data into the same \( d_1 \)-dimensional feature space as the vectors \( s_j \). For each observation \( (h_i, l_i) \),

\[
 l_i \xrightarrow{\text{embed}} h_i : h_i = W \cdot DNN(l_i), \tag{3}
\]

here \( DNN(\cdot) \), a function mapping from \( R^n \) to \( R^{n_{output}} \), represents a deep neural network. In our experiment, we empirically found that a 3-hidden-layer fully connected neural network using tanh as the activation function worked the best. The number of neurons in each hidden layer was 256, 256, 64. \( W \), a \( d_1 \)-by-\( n_{output} \) projection matrix, is used for modulating the data range and mapping the DNN’s output layer to the same high-dimensional feature space with \( s_j \), and \( n_{output} \) is the dimension of output layer. We will include more discussion about the performance of the neural network in the experimental section.

With these embeddings for each species and the environmental features at each observation site, we use the inner-product \( s_j^T h_i \) to score the relationship between \( \text{species}_j \) and environmental features in the observation site \( l_i \). Similarly, we use the inner-product \( \lambda_j^T \lambda_k \) to score the interaction between \( \text{species}_j \) and \( \text{species}_k \).

In order to simplify the presentation, we concatenate the vectors \( s_j \) and \( \lambda_j \) as the columns into two matrices.

\[
 S = (s_1, s_2, ..., s_n) \in R^{d_1 \times n}, \\
 \Lambda = (\lambda_1, \lambda_2, ..., \lambda_n) \in R^{d_2 \times n} \tag{4}
\]

Using the notations in equation(2), (3) and (4), we can formulate our DMSE model as follows,

\[
 \Pr(b_{ij} = 1 | l_i) = \Pr(r_{i,j} > 0), \\
 r_i \sim N_n(\mu_i, \Sigma), \tag{5}
\]

where \( \mu_i = S^T h_i = S^T(W \cdot DNN(l_i)) \) and \( \Sigma = \Lambda^T \Lambda \).

Here \( \mu_{i,j} = s_j^T h_i \) scores the habitat suitability of \( \text{species}_j \) at observation site \( l_i \) and \( \Sigma_{j,k} = \lambda_j^T \lambda_k \) represents the correlation between \( \text{species}_j \) and \( \text{species}_k \). According to the definition of multivariate normal distribution, we derive that

\[
 \Pr(b_{ij} | l_i) = \int_{L_1}^{R_1} \cdots \int_{L_n}^{R_n} f(x)dx_1...dx_n \tag{6}
\]

where

\[
 f(x) = \frac{1}{\sqrt{(2\pi)^n |\Sigma|}} \exp\left(-\frac{1}{2}(x - \mu)^T \Sigma^{-1} (x - \mu)\right),
\]

and \( L_j = \begin{cases} 0 & \text{if } b_{ij} = 1 \\ -\infty & \text{if } b_{ij} = 0 \end{cases} \) \( R_j = \begin{cases} +\infty & \text{if } b_{ij} = 1 \\ 0 & \text{if } b_{ij} = 0 \end{cases} \)

2.3 Training and Testing

We train our model by maximizing the log-likelihood on the observational data. The parameters that should be trained are the matrix \( S, \Lambda, W \) and the parameters in the deep neural network denoted by \( \theta_{DNN} \):

\[
 S, \Lambda, W, \theta_{DNN} = \arg\max_{S, \Lambda, W, \theta_{DNN}} \sum_{i=1}^{N} \log Pr(b_{i} | l_i) \tag{7}
\]

We use the stochastic gradient descent algorithm as proposed in [Duchi et al., 2011; Bottou, 2004] to optimize the log-likelihood function in equation (7).

In order to train and test our DMSE model, we need to be able to compute the integration in equation (6) and its derivatives with respect to each parameter. For the integration, we use an adaptive algorithm proposed in [Genz, 1992], which can calculate the cumulative distribution function (CDF) on multivariate normal distribution with a high accuracy (relative error \(< 10^{-6} \)).

To compute the derivative of \( \Pr(b_{ij} | l_i) \), one key observation is that if we can compute the derivative of \( \Pr(b_{ij} | l_i) \) with respect to \( \mu \) and \( \Sigma \), we can easily obtain other derivatives.
we want by simply applying the chain rule. Since the multivariate normal distribution is uniformly continuous, we first transform the derivative of the integral into the derivative of the density function as follows:

\[
\frac{\partial \log \Pr(b_i|l_i)}{\partial \mu} = \frac{1}{\Pr(b_i|l_i)} \int_{L_1}^{R_1} \ldots \int_{L_n}^{R_n} \frac{\partial f(x)}{\partial \mu} dx_1 \ldots dx_n
\]

\[
\frac{\partial \log \Pr(b_i|l_i)}{\partial \Sigma} = \frac{1}{\Pr(b_i|l_i)} \int_{L_1}^{R_1} \ldots \int_{L_n}^{R_n} \frac{\partial f(x)}{\partial \Sigma} dx_1 \ldots dx_n
\]

Using the definition of multivariate normal distribution, we derive the following equations:

\[
\frac{\partial f(x)}{\partial \mu} = f(x) \cdot F(\Sigma, \mu, x), \quad \frac{\partial f(x)}{\partial \Sigma} = f(x) \cdot G(\Sigma, \mu, x)
\]

where \( F(\Sigma, \mu, x) = \Sigma^{-1} (x - \mu) \),

\[
G(\Sigma, \mu, x) = -\frac{1}{2} (\Sigma^{-1} - \Sigma^{-1} (x - \mu) (x - \mu)^T \Sigma^{-1})
\]

According to equation (6), we know that

\[
\int_{L_1}^{R_1} \ldots \int_{L_n}^{R_n} \frac{f(x)}{\Pr(b_i|l_i)} dx_1 \ldots dx_n = 1
\]

Thus, we can consider \( \frac{f(x)}{\Pr(b_i|l_i)} \) as the density function of a distribution over a hyper-cube \( Q \subseteq \mathbb{R}^n \) corresponding to the integration range of equation(10). Thus, we can employ the Markov Chain Monte Carlo sampling method to estimate the derivative of \( \log \Pr(b_i|l_i) \) with respect to \( \mu \) and \( \Sigma \) as follows:

\[
\frac{\partial \log \Pr(b_i|l_i)}{\partial \mu} = \int_{L_1}^{R_1} \ldots \int_{L_n}^{R_n} \frac{f(x)}{\Pr(b_i|l_i)} F(\Sigma, \mu, x) dx_1 \ldots dx_n = E[F(\Sigma, \mu, x)]_{x \in Q} \approx \frac{1}{M} \sum_{k=1}^{M} F(\Sigma, \mu, x_k)
\]

\[
\frac{\partial \log \Pr(b_i|l_i)}{\partial \Sigma} = \int_{L_1}^{R_1} \ldots \int_{L_n}^{R_n} \frac{f(x)}{\Pr(b_i|l_i)} G(\Sigma, \mu, x) dx_1 \ldots dx_n = E[G(\Sigma, \mu, x)]_{x \in Q} \approx \frac{1}{M} \sum_{k=1}^{M} G(\Sigma, \mu, x_k)
\]

To make our model more efficient, we apply an enhancement for our model. Using the property of normal distributions, we know that \( \Pr(|r_{i,j} - \mu_j| > k \Sigma_{j,j}) < \frac{e^{-k^2/2}}{k \sqrt{2\pi}} \). As the result, we can make a cut-off on \( L_i \) and \( R_i \) which significantly reduces our sample range and increases the convergence rate in our sampling process.

3 Related Works

We refer the reader to [Elith and Leathwick, 2009] for a survey of general techniques used in species distribution modeling. Modeling approaches in this area vary depending on the type of observational data and application objectives. The most commonly available observational data records only where species have been detected and identified, known as presence-only data. The authors of [Phillips et al., 2004] developed the popular MaxEnt model using maximum entropy to estimate the population intensity. More recently, the connections between Poisson point processes and MaxEnt have been used to develop presence-only data models [Fithian and Hastie, 2013]. Other data collection protocols, like eBird, record both when species are and are not detected. These presence-absence datasets are typically modeled using a variety of statistical and machine learning methods including additive logistic regression, random forests, and boosted regression trees. Occupancy models [MacKenzie, 2006] account for imperfect detection of species by explicitly modeling hierarchically linked observation and occupancy processes, resulting in stronger ecological inferences [Guillera-Arroita et al., 2015; Hutchinson et al., 2011]. Species distribution models have also been extended to capture population dynamics using cascading models [Sheldon and Dietterich, 2011], Brownian Bridges [Horne et al., 2007], circuit theory [McRae et al., 2008], and non-stationary predictor response relationships [Fink et al., 2013]. Recent extensions to joint species distribution models focus on modelling the unobserved environmental factors which potentially drive the correlated distribution [Harris, 2015], and for spatiotemporal dynamics based on Gaussian processes [Thorson et al., 2016]. In machine learning literature, multi-label classification [Yu et al., 2005; Ji and Ye, 2009; Zhang and Zhou, 2010; Read et al., 2015; Jones et al., 2011; Jun et al., 2011] is also related to our work and can be applied to multi-species modeling. Most research in multi-label classification is based on ensemble of classifier chains (ECC), which is different from our approach and cannot provide direct information about the species correlation matrix. Among these previous work, [Harris, 2015], which also uses the latent random variables to model correlations, is most closely related to ours. However, their model can only handle a few (no more than 10) random variables to infer the unobserved factors which potentially drive the correlated distribution, and it ignores the interaction between species. In contrast, our DMSE method can handle hundreds of latent random variables for each species and can quantitatively measure the interaction among species. In the experiments section, we show that our DMSE model outperforms many aforementioned models.

Our model is also inspired by embedding methods which are widely applied to many areas, including music [Chen et al., 2012], language [Bengio et al., 2003; Mikolov and Dean, 2013], online purchase behavior [Rendle et al., 2010] etc. The core idea is to learn a vector (or other structure) to represent each of the data points, so that the interaction in the vector space reflects the semantic meaning in the original data. Embedding methods have been proven to have better generalization performance and to provide a better data visualization as well. [Rudolph et al., 2016] presents an embedding model that assumes an exponential family of conditional distributions, similar to Generalized Linear Models [McCullagh, 1984], to link observed quantities to latent embeddings that capture the semantic relationships of interest. Our DMSE model was developed independently of [Rudolph et al., 2016]\(^1\). While the probit model used in DMSE is in

\(^1\)We thank Liping Liu and David M. Blei for bringing up to our attention and discussing the Exponential Family Embedding in personal communications.
the exponential family, DMSE differs fundamentally from the work in [Rudolph et al., 2016]: The DMSE framework considers two heterogeneous contextual information feature sets (environmental features and interspecies relationships), it uses a deep neural network at the latent quantity level to extract high-level feature from environmental covariates and it couples the environmental and species embeddings into a predictive multi-species distribution model. It would be interesting to adapt the embeddings proposed in [Rudolph et al., 2016] and incorporate them into our DMSE setting. To our knowledge, we are the first ones to apply embedding methods with deep neural network structure to multi-species modelling.

4 Experiments

We work with crowd-sourced bird observation data collected from the successful citizen science project eBird [Munson et al., 2012]. One record in this dataset is often referred to as a checklist in which the bird observer reports all the species he/she detects as well as the time and geographical location of the observation site. Crossed with the National Land Cover Database for the U.S. [Homer et al., 2015], we can estimate the landscape composition of each observation site \( l_i \) with 15 different land types such as the percentage of the water, forest, grass, etc. For the use of training and testing, we transform all this data into the form \( (b_i, l_i) \) as described in the first paragraph of the Multi-Species Modelling section. The dataset for this experiment is formed by picking all the observation checklists from the Bird Conservation Region [Committee and others, 2000] (BCR) 13 in the last two weeks of May from the 2002-2012 which contains 39154 observations. May is a migration period for BCR 13, therefore a lot of non-native birds pass over this region, which gives us excellent opportunities to observe their habitat choice during the migration. Here we choose the top 100 most frequently observed birds as the species collection which covers 97.6% of the records in our dataset. In the experiments, we use a 5-fold cross validation to validate the multiple choices of hyper-parameters as well as evaluate the stability of models and we observe no overfitting between the loss on the validation vs test set during cross-validation.

4.1 What do embeddings look like?

We start by giving a qualitative impression of the embeddings produced by our method and visualized by t-SNE algorithm [Maaten and Hinton, 2008], which is a popular visualization method that can visualizes high-dimensional data by giving each datapoint a location in a two-dimensional map while, to a large extent, maintaining their original proximity. Fig.4 visualizes the embeddings of environmental preference and interactive behavior \((s_j, \lambda_j)\) of each species. In the picture, we manually assign the species into four main categories according to their habitat preference:

1. **Birds living near residential areas**, such as House Sparrow, Common Grackle, American Robin, Blue Jay, Mallard, Canada Goose, etc. Most of them are easy to find in the backyards, city parks, parking lots and agricultural fields. The presence ratio of these species are more than 25% of the records since they are easy for bird-watchers to find.

2. **Birds living in wetlands**, such as Swamp Sparrow, Northern Rough-winged Swallow, Killdeer, etc. that live near the water but mainly feed on insects.

3. **Water birds**, such as gulls, herons and cormorants which need a large amount of open water.

4. **Birds living in forest and pasture**, such as warblers, woodpeckers, nuthatches, thrushes, hawks, etc. These kinds of birds always live in the forest, grassland, pasture, shrubs, or near forest edges. These are the four categories that do not overlap with each other.

\( ^2 \)We get the habitat preference of birds from the website www.allaboutbirds.org
One can see in the left map of Fig. 4, the birds of the same category cluster tightly. For example, the birds living near the human settlements are all on the left, the birds living in wetlands and the water birds are on the right-top corner. Since the birds living in forest and pasture have a large habitat range, we further highlight three taxa in this category: the warblers, the woodpeckers and the raptors. It is interesting to note that the birds within a taxon have a high similarity of habitat preference which coincides with the field observation.

When it comes to the embeddings of correlation (the right map), it can be observed that in most cases, the species living in similar places have a relative higher correlation. However, one can find some interesting cases comparing the left map and the right map. For example, although the Mallard and Canada Goose are more common to see near human habitation, the occurrence of these two birds still has a high correlation with other water birds. What is more, in the left map, we find that the locations of Blue Jay and American Robin are not very close, but from the right map, we know that they have a very high correlation which coincides with the ecological relationship as we described in the introduction section.

### 4.2 Predictive Performance of DMSE

While the visualization provides interesting qualitative insights, we now provide a quantitative evaluation of the model quality based on predictive power. In our experiment, we analyze the performance of DMSE for modeling both single-species and multi-species. Here we use two metrics to analyze the predictive performance: (1) Area under Curve (AUC), i.e. the area under receiver operating characteristic (ROC) curve which is used to describe the ability of the model to rank outcomes, and (2) the log-likelihood, i.e. \( \sum_{i=1}^{N} \log P_r(b_i|l_i) \), which measures the calibration of the predicted probabilities of species occurrence.

#### DMSE’s performance on a single-species model

We compare the single-species predictive performance of DMSE with random forest (RF) model and SVM in terms of AUC. Random forest is one of the standard techniques used to model single species distribution in a wide variety of ecological and conservation applications (e.g. [Herrando et al., 2017; Elith and Leathwick, 2009]) and SVM is also a popular and robust model which has been successfully used for numerous applications. We implemented RFs and SVMs using python-sklearn. The number of trees in RFs was 1000, which saturated the predictive performance. The kernel of SVMs was RBFs, which perform well across a range of applications. Here we also analyze the effect of the deep neural network in the DMSE model by analyzing the performance of a DMSE model, in which we only use projection matrix \( W \) to embed the environmental features. We test these four models on different species from very common to rarely seen. As shown in Fig. 5, the deep neural network gives us a significant boost on the predictive power of DMSE. We expect a similar performance boost when we incorporate deep structures into other relevant models, such as the exponential family embedding model in [Rudolph et al., 2016]. With the help of deep neural network, our DMSE model outperforms other models.

**What are the effects of correlation?**

We now explore whether the correlation plays an important role in multi-species modelling. We start by comparing the performance of multi-species DMSE and the single version of DMSE in modelling two-species distribution. The single version of DMSE can be thought of as the original DMSE model but with an identity correlation matrix, which means we model the multi-species distribution by modelling the distribution of each species independently without their correlation. Here we use log-likelihood instead of AUC to analyze models’ performance because the AUC averaged across species still values the distribution of each species separately, which does not fully reflect the benefit of modeling correlation. According to our experimental results, the multi-species DMSE outperforms the single version on all the species pairs that we have tried. Because of space limitation, we only show the performance on 3 pairs of species.

As shown in Fig. 6, multi-species DMSE has a substantial improvement compared with the single version of DMSE, which reflects the important role of inter-species correlation. Furthermore, we provide Table. 1, which quantitatively measures the interaction between some species pairs with relatively high correlation. These pairs have been validated by domain experts.

In addition, we compare our DMSE model with (1) the single version of DMSE, (2) the ensemble of classifier chains (ECC) method [Jones et al., 2011] which is a standard-class of models used for multi-label prediction within the discipline of machine learning (We use single version of DMSE as building blocks of the classifier chain) and (3) MISTNET model proposed in [Harris, 2015], a recent extension of species distribution model that models unobserved environmental factors which potentially drive the correlated distribution of multiple species.

During the training and testing, all these models use the same environmental feature as their input, which is the land-
Table 1: The list for species pairs with high correlation. The correlation here is derived from covariance matrix $\Sigma$.

<table>
<thead>
<tr>
<th>Species Name</th>
<th>Species Name</th>
<th>correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red-eyed Vireo</td>
<td>Eastern Wood Pewee</td>
<td>0.607</td>
</tr>
<tr>
<td>Common Grackle</td>
<td>Red-winged Blackbird</td>
<td>0.604</td>
</tr>
<tr>
<td>European Herring</td>
<td>Great Black-backed Gull</td>
<td>0.580</td>
</tr>
<tr>
<td>Yellow Warbler</td>
<td>Common Yellowthroat</td>
<td>0.567</td>
</tr>
<tr>
<td>Blue Jay</td>
<td>American Robin</td>
<td>0.535</td>
</tr>
<tr>
<td>Common Grackle</td>
<td>American Robin</td>
<td>0.510</td>
</tr>
<tr>
<td>Blue Jay</td>
<td>Northern Cardinal</td>
<td>0.504</td>
</tr>
<tr>
<td>American Crow</td>
<td>American Robin</td>
<td>0.493</td>
</tr>
<tr>
<td>Common Grackle</td>
<td>European Starling</td>
<td>0.475</td>
</tr>
<tr>
<td>European Starling</td>
<td>Red-winged Blackbird</td>
<td>0.474</td>
</tr>
</tbody>
</table>

Figure 7: As the number of species becomes larger, the performance of multi-species DMSE becomes better and better compared with single species DMSE and other models. This figure shows the performance difference of all models against the single version of DMSE.

6 Acknowledgements

We are thankful to the Cornell Lab of Ornithology for providing eBird data for this research. This research was supported by National Science Foundation (Grant Number 0832782, 1522054, 1059284, 1356308), ARO grant W911-NF-14-1-0498, the Leon Levy Foundation and the Wolf Creek Foundation.

References


[Guillera-Arroita et al., 2015] Gurutzeta Guillera-Arroita, José J Lahoz-Monfort, Jane Elith, Ascelin Gordon, Heini Kujala, Pia E Lentini, Michael A McCarthy, Reid Tingley, and Brendan A Wintle. Is my species distribution model fit for purpose? matching high-dimensional feature space via a deep neural network. Our DMSE model significantly outperforms existing models on multi-species distribution modelling. Additionally, we demonstrate the benefit of using a deep neural network for feature extraction and show how they improve the predictive performance of species distribution modelling. The ability to visualize the learned embeddings is also a key feature for easy interpretability and open-ended exploratory data analysis. Furthermore, the embedding models described in the paper can easily be adapted and extended to include further information (e.g., spatio-temporal information), providing many directions for future work.

5 Conclusion

We present a novel Deep Multi-Species Embedding model that can quantitatively capture inter-species correlations of hundreds of species simultaneously, by jointly embedding vectors corresponding to multiple species as well as vectors representing environmental covariates into a common

...
data and models to applications. *Global Ecology and Biogeography*, 2015.


