Template to prepare preprints and manuscripts using markdown and github actions

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Purpose: This template provides a series of scripts to render a markdown document into an interactive website and a series of PDFs.

Motivation: It makes collaborating on text with GitHub easier, and means that we never need to think about the output.

Internals: GitHub actions and a series of python scritpts. The markdown is handled with pandoc.

Keywords: pandoc pandoc-crossref github actions

Introduction

This document is for brainstorming the Colorado bumblebee metaweb prediction project.

1 Data

The data consists of two parts, the field data collected at three stations across Colorado, and the GBIF metaweb data aggregated and cleaned by Julian R. and a student.

2.1. 1.1 Field data This data consists of nearly-daily records of visits between bee and flower species for the months August-October. Data is collected from three field sites: 1) Rocky Mountain Biological Lab (RMBL) 2) Niwot Ridge 3) Pikes Peak

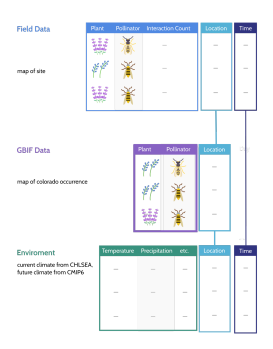
each with tens of plots each across an elevational gradient, collected for 7/4/3 summers total.

To be determined: specific number of plots per site and years per site

2.2. 1.2 GBIF metaweb This data of a network of bee/flower species which have a record of both occurring on GBIF for locations at some spatial grain

To be determined: how does the scale of spatial aggregation affect properties of metaweb, how much taxonomic overlap is there with field data?—We have a lot of field data on interactions, but we can't validate predictions on taxa that aren't present in field data. As a result validation can only be done on interactions for which both species appear in field data.

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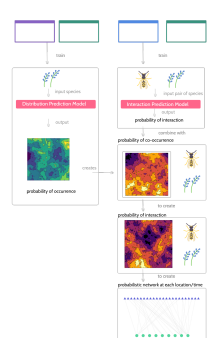


Figure 1 todo

3

Methods

Where X_{ij} is and L_{abc} is .

as in gravel et al 2017

$$P(X_{iy}, X_{jy}, L_{ijy}|E_y) = P(X_{iy}, X_{jy}P(L_{ijy}|X_{iy}, X_{jy}, E_y)$$

Then decompose probaility of co-occurence as

$$P(X_{iv}, X_{iv}) = P(X_{iv})P(X_{iv})$$

3.1. A predictive model to make spatially explicit network prediction The goal is two have two predictive models: interaction-predictor model and a distribution-predictor model (a la Strydom & Catchen et al. 2021, figure 2).

The interaction-predictor model, $f_i(s_i, s_j, \theta_i)$ predicts interaction based on species-level features (s_i, s_j) , and is trained on the field-data.

These features could include Phylogeny (to be determined: how available are genomes or trees for these species) Environment/Climate Traits (to be determined: what trait data is available, how annoying is it to clean) Time (only for the phenology model, see 3.2 and 3.3)

The distribution-predictor model, $f_s(s_i, \vec{x}, t)$ is trained on GBIF data to predict the occurrence of species with features si at a location in space x, and time t. Many options here. Here the species level features could be Climatic variables derived from remote sensing products. Co-occurrence to make a JSDM Potentially weighted by phenology information from field data. Time (only for the phenology model, see 3.2 and 3.3)

3.2. Combining distribution-predictor and interaction-predictor models Can split this into two based on how the distribution-predictor works. If f_s predicts co-occurrence, then draw the species pool first and predict interactions between the species in that pool. If f_s is a single-species SDM, get the occurrence probability for each species p_s and compute the probability of observing interaction as function of the product of occ. prob.

4
Results
After comparing different combinations of features/model structures and finding the 'best' performing model on validation data.
4.1. Figure one: spatial species pool and network prediction Figure that is two panels: a map of total species richness and a map of network properties across Colorado. This model doesn't consider time, only other predictors.
4.2. Figure two: Phenology Same as figure one but consists of maps but at different times of the year (e.g. March, June, August) and uses both an interaction-predictor and distribution-predictor that incorporate time into predictions
4.3. Figure three: Climate Much as climate change has shifted temperature gradients to get warmer toward the poles, it has also moved temperature gradients up in elevation.
We can get a CMIP6 forecast of temperature and precipitation, and then predict how many observed interactions in the field data will no longer have their composing species' distributions overlap. Decompose temporal component of overlap from spatial component.
5
Discussion
5.1. Acknowledgements
6
References