Phylogenetic Transfer Learning
Michael D. Catchen ^{1,2}
¹ McGill University ² Québec Centre for Biodiversity Sciences
Correspondance to:
Michael D. Catchen — michael.catchen@mcgill.ca

Substance: Find a bunch of ASR datasets, do crossvalidation on tip states, test predictive efficacy on inputing 'hidden' species.

Premise: Imputing traits among extant species using ancestral state resconstruction (ASR) can be useful for a wide variety of purposes in ecology

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What is Phylogenetic Transfer Learning

Ancestral State Reconstruction

- 3 Ancestral State Reconstruction (ASR; also called ancestral character reconstruction, character estimation, or
- 4 character mapping) is a core topic in phylogenetics (**Joy2016AncRec?**).
- The goal is, given an estimate of a phylogeny \mathcal{P} and trait values T_i for each extant species in the phylogeny i, to
- 6 estimate the value of that trait at some historical point in the phylogeny (typically at the node representing the
- 7 MRCA of a clade of interest).
- ⁸ To do this, one assumes a statistical model of evolution (and often tries to infer the best among a set of candidate
- 9 models). Depending on whether the trait of interest is discrete or continuous, the models are typically
- discrete-space Markov chains (where the transition matrix is a target of inference) or, in the continuous setting,
- either Brownian Motion or an Ornstein-Uhlenbeck (OU) process, where the former is used for neutral evolution
- and the latter is used when there is hypothesized selection on the trait.
- The methodology used to fit models has followed the historical progression from maximum parsimony models
- 14 (a naive approach that favors as few evolutionary changes as possible in discrete traits) to maximum likelihood
- estimation. Modern methods revolve on using Bayesian methods, which has the direct benefit of including
- uncertainty estimates in infered ancestral states.

17 Phylogenetic Transfer Learning

- The core insight of Phylogenetic Transfer Learning (PTL) is to take the the model of evolution and ancestral
- state infered via ASR, and simulate the parameterized evolutionary model forward from the ancestral node to
- get an estimate of extant trait values (with uncertainty) for species without trait observations T_i .
- 21 The transfer learning component in particular comes from the first use of this methods to inpute latent
- representations of species (Strydom2022FooWeb?; Strydom2022GraEmb?)

Substance of the paper

Simulated phylogenies

- Various properties of radiation
- 26 2. Real data

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- ²⁷ Relevant questions: What phylogenetic scales can PTL give robust predictions There is a clear upper limit (if
- 28 the MRCA for set of species is too long ago, there will be massive variance in predicted state at the tip) There
- is also a lower limit (if there is HZT and the phylogeny is not a reliable proxy for how traits are evolving) -
- Robustness to noise is trait measurements? How does this very w/ amount and different types of selection

Possible Applications

- SDMs: mechanistic niche predictions on subsets of species
- Connectivity (Catchen2023ImpEco?)