# Should an individual based mechanism be implemented in the protracted birth death model?

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#### 1. Abstract

Speciation models are used to construct phylogenetic trees. Most of the currently used models, including the protracted birth-death model, lack a mechanism on individual level. The models assume several constant factors to determine if a new lineage should arise or an old lineage should go extinct. An individual based model might give us more realistic phylogenies and answer evolutionary questions better than the currently used (protracted) birth-death model

# 2. Introduction

#### 2.1 Speciation models

#### 2.1.1 The need for speciation models

Speciation models help us construct phylogenies to answer evolutionary biological questions. [REF 4] They are used to give a better understanding of speciation in nature.

#### 2.1.2 Birth-death model

The standard birth-death model [5] shows an accelerating increase in the number of species. This is not shown in phylogenetic trees, which have a slowdown in the increase of number of lineages. [2] The birth-death model is one of the simplest of models [4] that has a constant speciation rate  $(\lambda)$  and a constant extinction rate  $(\mu)$ .

#### 2.1.3 Protracted birth-death model

By assuming that speciation is not instant and happens over a longer period of time we can change the birth-death model to a protracted birth-death model. "Good" species have a chance of making incipient species with rate  $b_g$  that will become good species with rate  $\lambda$ . Incipient can initiate speciation with rate  $b_i$ . Both good and incipient species can go extinct with rate  $\mu_g$  and  $\mu_i$  respectively. [2] [REF 4]

#### 2.1.4 Relationship between birth-death and protracted birth-death

# 2.1.5 Other speciation models

[2] [REF]

# 2.1.6 Problems of speciation models

# 2.2 The use of mechanisms to model speciation

# 2.2.1 Ecological and sexual selection model

[1]

# 2.2.2 Bateson-Dobzhansky-Müller model

[REF Jorik]

# 2.3 Differences protracted birth-death model and ecological and sexual selection model

[1][2]

#### 2.4 This study

This study compares the phylogenetic trees of the protracted birth-death model and ecological and sexual selection mechanism, by changing parameters to get the resulting trees as similar as possible.

- 2.5 Novelty
- 2.6 Analysis

[3]

# 2.7 Expectations

# 3. Methods

#### 3.1 Parameters

#### 3.1.1 PBD

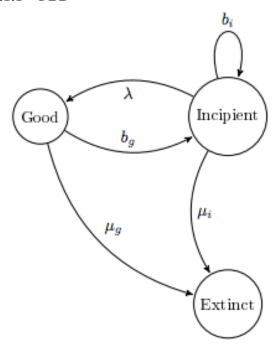


Figure 2: The states and transitions of a species.  $b_i$ : speciation-initiation rate of incipient species.  $b_g$ : speciation-initiation rate of good species.  $\lambda$ : speciation completion rate.  $\mu_i$ : extinction rate of incipient species.  $\mu_g$ : extinction rate of good species. Figure after Etienne et al, 2014, Evolution

[2]

# 3.1.2 Ecological and sexual selection model

[1]

# 3.2 Creating species trees

# 3.3 Analysis

#### 3.3.1 nLTT

[3]

# 3.3.2 γ

The Gamma statistic (Pybus & Harvey 2000) is given by:

Pybus, O. & Harvey, P. (2000) Testing macro-evolutionary models using incom-

plete molecular phylogenies. Proceedings of the Royal Society B: Biological Sci- ences, 267, 2267–72.

#### 3.4 Peripherals

#### 3.4.1 Software

https://github.com/Lumphie/Research\_phylogenies

#### 3.4.2 Hardware

# 4. Results

# 5. Discussion

# References

- [1] Van Doorn, G. S., & Weissing, F. J. (2001). "Ecological versus sexual selection models of sympatric speciation." Selection, 2, 17 40.
- [2] Etienne, Rampal S., and James Rosindell. (2011) "Prolonging the past counteracts the pull of the present: protracted speciation can explain observed slowdowns in diversification." Systematic Biology: syr091.
- [3] Janzen, Thijs, Höhna, Sebastian, Etienne, Rampal S. (2015) "Approximate Bayesia n Computation of diversificatio nrates from molecular phylogenies: introducing a newefficient summary statistic, the nLTT." Methods in ECology and Evolution 6, 566 575.
- [4] (My current work)
- [5] Kendall, D. G. (1948) "On some modes of population growth giving rise to R.A. Fisher's logarithmic series distribution." Biometrika, 35, 6 15.