

# Ngesh: a Python library for synthetic phylogenetic data

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#### Software

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### Summary

This work presents ngesh, a Python library for simulating phylogenetic trees and data, designed for usage in development, debugging, and benchmarking of analysis pipelines and methods for phylogenetic inference, particularly in historical linguistics and stemmatics. The package generates reproducible stochastic simulations of evolution according to various criteria, including character mutation rates and probability of horizontal transfer, and its results can include the simulation of inadequate data compilation and sampling. Different output formats are supported, both for visualization (such as plain text and with integrated graphical viewers) and for software interoperability (such as Newick and NEXUS).

## **Background**

Computational phylogenetics is being increasingly accepted in fields beyond biology, such as historical linguistics (Bouckaert et al., 2012) and stemmatics (Robinson, 2016). Stochastic simulations, long advocated for natural sciences in general (Bailey, 1964) and genetics in specific (Foote et al., 1999; Harmon, 2018), are not used enough in these fields. Nonetheless, they are very desirable, allowing to evaluate evolutionary analogies, models, and performance through vast amounts of simulated histories, without limits imposed by data availability and collection time, with quantifiable precision of results. Simulations can also be used to perform fuzzy testing of software and to support studies on which evolutionary models, processes, and evolutionary parameters better match the observed phenomena.

The ngesh library is a tool that allows to perform such simulations, designed for easy integration into phylogenetic pipelines. It can generate reproducible trees and correlated data following both user-established parameters, such as ratios of birth and death, and constraints, such as branch lengths and minimum number of taxa. The library can label taxa progressive enumeration or with random names that are easy to pronounce (e.g. "Sume" and "Fekobir") or which imitate the binominal nomenclature (e.g. "Sburas wioris" and "Zurbata pusso"). Character evolution related to the tree topology can likewise be simulated, including *ex novo* mutations and horizontal gene transfers. Results can be manipulated in diverse manners, for example by pruning extinct leaves or simulating uneven sampling. The simulated trees are standard ETE3 objects (Huerta-Cepas et al., 2016) and may be exported into different formats such as Newick trees, ASCII-art representation, and tabular lists.

#### Statement of need

The library addresses the need of more tools to investigate and teach phylogenetics in historical linguistics and stemmatics. As a building block for evaluating pipelines of analysis, it is an



alternative to the basic technique of randomizing taxa placement in existing cladograms, and to simpler tools such as the one by Noutahi (2017) or the populate() method of ETE3's Tree class (Huerta-Cepas et al., 2016). While there are many other alternatives available for simulating trees, including TreeSim (Stadler, 2011), geiger (Pennell et al., 2014), ape (Paradis & Schliep, 2018), and DendroPy (Sukumaran & Holder, 2021), ngesh compares favorably in historical linguistics and stemmatics. For the former, it provides default parameters that produce trees closer to those found in the field, particularly in terms of the simulation of horizontal transfers (i.e., loanword), all while using formats that better fit the existing linguistic pipelines, such as CLDF (Forkel et al., 2018), and laying ground for the usage of different character values (such as sound changes) besides the default cognate-sets for modelling lexical replacement. For the latter, where Bayesian phylogenetics have been gaining traction at a slower pace, the library constitutes the first general-purpose tool available and should help make these methods for popular.

## Installation, Usage, & Examples

Users can install the library with the standard pip tool for managing Python packages. Trees can be generated from the command-line, defaulting to small phylogenies in Newick format:

```
$ ngesh
(Ukis:1.11985,(Koge:0.880823,(Rozkob:0.789548,(Meu:0.706601,
(((Felbuh:0.189693,Kefa:0.189693)1:0.117347,((Epib:0.153782,
Vugog:0.153782)1:0.0884745,Puluk:0.242256)1:0.0647836)1:0.0469885,
Efam:0.354028)1:0.352573)1:0.0829465)1:0.0912757)1:0.23903);
```

The tool supports both configuration files and command-line flags that take precedence over the former. Here we specify a model to generate Nexus data for a reproducible Yule tree, with a birth rate of 0.75, at least 5 leaves, "human" labels, and 20 presence/absence features:

```
$ cat my_tree.conf
[Config]
labels=human
birth=0.75
death=0.0
output=nexus
min_leaves=5
num chars=20
$ ngesh -c my_tree.conf --seed 12345
begin data;
  dimensions ntax=6 nchar=33;
  format datatype=standard missing=? gap=-;
  matrix
          111110110111011011010101000100110
R1172
          1111110101101111101100010010011001
Lenlar
Mukom
          111110111011011011101001000100110
Pagil
          111110110111011011100100100100110
          111110110111011011100011001001010
Suglu
Wite
          111110110111011011100101000100110
end;
```

Despite the benefit of a stand-alone tool, the package is designed to be run as a library. The two primary functions are gen\_tree(), which returns a random tree, and add\_characters(),



which adds character evolution data to a tree. Users can generate random trees without character information or simulate character evolution within existing trees, including non-simulated ones.

Besides the write() method above, which outputs Newick trees, results can be exported in either NEXUS format with tree2nexus() or in a textual tabular format with tree2wo rdlist(). Phylogenetic reconstruction can then be carried either by manually building an XML model for BEAST2 (Bouckaert et al., 2019) (normally with the aid of the graphical interface BEAUTi) or by using tools such as BEASTling (Maurits et al., 2017), producing a tree distribution. This distribution can be summarized to a maximum clade credibility ("MCC") tree with phylogenetic packages, allowing both visual and quantitative comparisons. A demonstration of such steps is provided with the user documentation ("Integrating with other software").

# **Code and Documentation Availability**

The ngesh source code is available on GitHub at https://github.com/tresoldi/ngesh. User documentation is available at https://ngesh.readthedocs.io/.

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