

# User manual for pyvolve v1.0

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## 1 Introduction

Pyvolve (pronounced “pie-volve”) is an open-source python module for simulating genetic data along a phylogeny according to Markov models of sequence evolution. The module is available for download on [github](#) (and see [here](#) for API documentation). Note that pyvolve has several dependencies, including [BioPython](#), [NumPy](#), and [SciPy](#). These modules must be properly installed and in your python path for pyvolve to work properly. Please file any and all bug reports on the github repository [Issues](#) section.

Pyvolve is written such that it can be seamlessly incorporated into your python pipelines without having to interface with external software platforms. However, please note that for extremely large (>1000 taxa) and/or extremely heterogeneous simulations (e.g. where each site evolves according to a unique evolutionary model), pyvolve may be quite slow and thus may take several minutes to run. Faster sequence simulators you may find useful include (but are certainly not limited to!) [Indelible](#) [1] and [indel-Seq-Gen](#) [8].

Pyvolve supports a variety of evolutionary models, including the following:

- Nucleotide Models
  - Generalized time-reversible model [9] and all nested variants
- Amino-acid exchangeability models
  - JTT [4], WAG [10], and LG [6]

- Codon models
  - Mechanistic ( $dN/dS$ ) models (MG-style [7] and GY-style [2])
  - Empirical codon model [5]
- Mutation-selection models
  - Halpern-Bruno model [3], implemented for codons and nucleotides

Note that it is also possible to specify custom matrices (detailed in section 7 below). Both site-wise and temporal (branch) heterogeneity are supported. Sequences are simulated according to standard methods [11].

## 2 Basic Usage

Similar to other simulation platforms, pyvolve evolves sequences in groups of **partitions**. Each partition has an associated size and model (or set of models, if branch heterogeneity is desired). All partitions will evolve according to the same phylogeny; if you wish to have each partition evolve according to a distinct phylogeny, I recommend performing several simulations and then merging the resulting alignments in the post-processing stage.

Pseudocode for a simple simulation is given below.

```

1  # Import the pyvolve module
2  import pyvolve
3
4  # Read in tree along which pyvolve should simulate
5  my_tree = pyvolve.read_tree(file = 'file_with_tree_for_simulating.tre')
6
7  # Define and construct evolutionary models
8  my_model = pyvolve.Model(<model_type>, <custom_model_parameters>)
9  my_model.construct_model()
10
11 # Define partitions
12 my_partition = pyvolve.Partition(models = my_model, size = 100)
13
14 # Evolve partitions with the callable Evolver() class
15 pyvolve.Evolver(tree = my_tree, partitions = my_partition)()
```

By default, sequences will be output to a fasta-formatted file called “simulated\_alignment.fasta”. Two additional tab-delimited files, called “site\_rates.txt” and “site\_rates.info.txt” are also output. These files provide useful information when heterogeneity (either site or branch) is implemented. The former file indicates to which partition and rate category (if no rate heterogeneity specified, these values will all be 1) each site belongs, and the latter file provides more specific information about each rate category, in particular its associated partition, probability, and value. To suppress creation of either file, or to change the file name, use the respective arguments `ratefile` and `infofile` when initializing an `Evolver` instance. Providing a value of either `None` or `False` will suppress file creation, or simply provide a string giving the desired output file name. To change the outputted alignment file name, set the argument `seqfile` (similar, providing `False/None` will suppress file creation). The sequence file format can also be changed with the argument `seqfmt`. Biopython is used to write sequence files, so consult the Biopython manual for available formats.

`<model_type>` is the type of model matrix. `<custom_model_parameters>` is a *dictionary* of parameters for your chosen model. See below for available model types and associated parameter keys.

### 2.1 Nucleotide Models

```

1  # Import the pyvolve module
2  import pyvolve
3
4  # Read in tree along which pyvolve should simulate
```

```

5 my_tree = pyvolve.read_tree(file = 'file_with_tree_for_simulating.tre')
6
7 # Define and construct evolutionary models
8 nucleotide_rates = {'AC':0.5, 'AG':0.2, 'AT':0.3, 'CG':1.5, 'CT':1.6, 'GT':3.4} # symmetric. all unspecified
   are set to 1.
9 my_model = pyvolve.Model('nucleotide', nucleotide_rates)
10 my_model.construct_model()
11
12 # Define partitions
13 my_partition = pyvolve.Partition(models = my_model, size = 100)
14
15 # Evolve partitions with the callable Evolver() class
16 pyvolve.Evolver(tree = my_tree, partitions = my_partition)()

```

## 2.2 Amino-acid models

## 2.3 Mechanistic codon models

## 2.4 Empirical codon models

## 2.5 Mutation-selection models

# 3 Site-wise heterogeneity

## 3.1 Nucleotide and amino-acid models

## 3.2 Codon models

## 3.3 Mutation-selection models

# 4 Temporal heterogeneity

# 5 Building a vector of stationary frequencies

# 6 Matrix scaling options

# 7 Using custom rate matrices

This is my first python example:

```

1 from pyvolve import *
2
3 # Read in a newick tree
4 t = read_tree(file = "myfile.tre")
5
6 # Construct state frequency vector. Optional!
7 f = EqualFrequencies("amino")
8 freqs = f.construct_frequencies(type = "codon")
9
10 # Build the evolutionary model
11 m = Model("GY94", {'state_freqs':freqs, 'omega':1.5, kappa:3.4})
12 m.construct_model()
13
14 # Initialize partitions
15 p = Partition(models = m, size = 100)
16
17 # Evolve, and call.
18 Evolver(partitions = p, tree = t, seqfile = "sequences.phy", seqfmt = "phylip")()

```

## References

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