Phylogenetics

Phytest: quality control for phylogenetic analyses

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Abstract

Motivation: The ability to automatically conduct quality control checks on phylogenetic analyses is becoming more important with the increase in genetic sequencing and the use of real-time pipelines e.g. in the SARS-CoV-2 era. Implementations of real-time phylogenetic analyses require automated testing to make sure that problems in the data are caught automatically within analysis pipelines and in a timely manner. Here, we present Phytest (version 1.1) a tool for automating quality control checks on sequences, trees and metadata during phylogenetic analyses.

Results: Phytest is a phylogenetic analysis testing program that easily integrates into existing phylogenetic pipelines. We demonstrate the utility of Phytest with real-world examples.

Availability and implementation: Phytest source code is available on GitHub (https://github.com/phytest-devs/phytest) and can be installed via PyPI with the command ‘pip install phytest’. Extensive documentation can be found at https://phytest-devs.github.io/phytest/.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Phylogenetics is increasingly performed in automated pipelines, run with increasing frequency as sequence data becomes more readily available e.g. during the SARS-CoV-2/Coronavirus disease 2019 (COVID-19) pandemic. The frequency of runs and complexity of these pipelines can result in the introduction of erroneous data causing failures, or worse, incorrect results and conclusions. Manually checking analyses for errors can be repetitive and laborious, wasting researchers time. While automated testing software exists for code, no such testing frameworks are available to easily test for errors in phylogenetic analyses.

Here, we present Phytest (version 1.1.0) a tool for automating quality control checks on sequence, phylogenetic trees and associated metadata files during phylogenetic analyses. Phytest ensures that phylogenetic analyses meet user-defined quality control standards. Phytest can be installed from the Python Package Index (PyPI) using the command ‘pip install phytest’.

2 Implementation and usage

Phytest is based on the popular Python testing framework Pytest (Krekel et al., 2004). It allows users to write tests for their phylogenetic analyses the same way they write tests for their code (Fig. 1). Phytest has been developed as a command-line interface (CLI), Python module and Pytest plugin, providing multiple methods of invocation. It provides many convenient helper functions for testing phylogenetic analyses including methods for testing sequences, alignments, trees and metadata files.

Phytest is easily extendable and provides a simple interface for writing custom phylogenetic tests. The interface follows the Pytest model of testing i.e. tests are defined as Python functions containing assert statements that are collected and evaluated at run-time. Tests that fail are captured and reported to the user allowing for repeatable and automated testing.

Phytest injects special fixtures into test functions, allowing for easy evaluation and testing of phylogenetic data structures. These fixtures provide the standard Biopython (sequences, alignments and trees) and Pandas (metadata) class methods as well as special assert methods for testing these data structures (Cock et al., 2009; McKinney et al., 2010). For example, the Phytest Sequence class implements the method Sequence.assert_percent_GC. Calling this method on the fixture object with the expected GC-content e.g. sequence.assert_percent_GC(38) will raise an error if the percent of G and C nucleotides in the sequence is not equal to 38%. Many methods also provide maximum and minimum arguments so the upper and lower bounds can be tested e.g. sequence.assert_percent_GC(min=30, max=40).

All Phytest assert methods also provide a warning flag, e.g. sequence.assert_percent_GC(38, warn=True) causing the method to raise a warning instead of an error if the test fails. In an automated pipeline, this provides a way to inform the user of potential problems without causing the pipeline to fail. The warning flag can be set automatically by calling the method with the warn_prefix instead of assert e.g. sequence.warn_percent_GC(38).

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fitting a molecular clock and estimating evolutionary rates and dates, such as when using BEAST (Bouckaert et al., 2019; Drummond and Rambaut, 2007). A repository containing the code for this example can be found at https://github.com/phytest-devs/phytest-temporal-signal-example. Here, we use data from the TempEst tutorial https://beast.community/tempest_tutorial. TempEst is a useful program for performing temporal signal analysis, however, it is not possible to easily automate the TempEst graphical user interface (Rambaut et al., 2016). Internally, Phytest uses TreeTime to perform a root-to-tip regression, allowing users to automate temporal signal testing (Sagulenko et al., 2018). The `tree.assert_root_to_tip` method is used for testing temporal signal and provides arguments for testing the coefficient of determination ($R^2$), regression slope (a crude estimate of the evolutionary rates) and root date.

**4 Conclusions**

Phytest is a flexible and powerful tool for reproducibility and automation. New data are often incorporated or manually edited when developing or optimizing a phylogenetic analysis pipeline. Phytest ensures that analyses meet user-specified standards each time they are run. We believe Phytest will be increasingly useful as the use of automated phylogenetic pipelines increases, especially in the field of real-time phylogenetic analysis.

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**Conflict of Interest:** None declared.

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