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Automagic phylogenies

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1 Introduction

This report provides an overview of a software program designed to generate phylogenetic trees. Phylogenetic trees are graphical representations of evolutionary relationships among species or groups of organisms. The program utilizes various algorithms and data inputs to generate accurate and comprehensive phylogenetic trees. This report will provide a brief overview of the features and capabilities of the software, as well as its intended use and target audience. The software program is designed to be user-friendly and accessible for both researchers and educators in the field of evolutionary biology. It integrates advanced algorithms for tree construction, allowing for the analysis of large and complex datasets. The program also includes visualization tools for tree presentation, as well as options for customizing and annotating the tree output. Additionally, the software can import and export data in a variety of formats, making it easy to integrate with other analysis tools. The program is intended to provide a comprehensive and efficient solution for phylogenetic tree construction and analysis, and is an essential tool for anyone studying evolutionary relationships among species or groups of organisms.

2 Background

In this section will be provided a quick background information on the field of phylogenetics and the challenges associated with creating phylogenetic trees.

Phylogenetics is a study that aims to understand the evolutionary relationships among vast groups of similar organisms. It uses molecular biology to achieve to compare the genetic and morphological characteristics of different organisms, infering their evolutionary relationships. The main goal of this process is to construct evolutionary/phylogenetic trees, which depict the evolutionary relationships among different organisms.

The process of creating a phylogenetic tree can be challenging. One of the main challenges is the availability of data. For example, it can be difficult to obtain high-quality genetic data for a certain group of organisms. The complexity of determining evolutionary relationships can be compounded by various factors such as, the method used, the type of data, and the assumptions made. Also, the construction of a phylogenetic tree assumes that similarities among organisms are the result of a shared ancestry, but it's possible that similarities may have developed independently within different groups of organisms.

Finally, creating a phylogenetic tree requires making choices about the appropriate model and the appropriate method for inferring relationships, like Maximum likelihood, Bayesian and Distance-based methods. Selecting the most fitting model can be challenging, but

fortunately, there are resources available to assist in making the best choice.

The field of phylogenetics requires expertise from multiple areas, like molecular biology and computer science. Phylogenetic trees can offer significant insights into the evolutionary connections among organisms, however, it is crucial to keep in mind the difficulties and ambiguities that can arise during the creation of these trees.

3 Methodology

The methodology for building phylogenetic trees involves several key steps, including data acquisition, data processing, and tree inference. Data acquisition involves obtaining high-quality genetic or morphological data for each of the organisms being analyzed. Data processing involves cleaning, organizing, and transforming the data into a format that is suitable for tree inference. Finally, tree inference involves using a variety of algorithms and models to construct the phylogenetic tree based on the processed data. In the next subsections it will be explained every part of the program from the data acquisition to the tree build and what methods were used.

3.1 Environment and WorkFlow Management

In order to ensure consistency and reproducibility of our program, we utilized Docker as our virtualization platform. Docker allowed us to package our software and dependencies into a single container, which could be run on any system with Docker installed. This eliminated the need for manual installation and configuration of dependencies, making our program easier to run and share.

To manage and automate the execution of our program, we used Snakemake, a workflow management system. Snakemake enabled us to specify the dependencies and rules for our pipeline, making it easy to run and scale our program. The Snakemake file served as a blueprint for the pipeline, defining the input, output, and steps required to complete the program. This allowed us to simplify and streamline the execution process, while still maintaining full control and visibility over the workflow.

3.2 Data Acquisition

For the data acquisition we used the EntrezAPI, which The Entrez API is a component of the NCBI (National Center for Biotechnology Information) programmatic access to the vast collections of data maintained by the NCBI. The API provides a set of programmatic tools for accessing NCBI databases, including PubMed, GenBank, and others. The API allows developers to retrieve and manipulate data in a format that is suitable for analysis and integration into other programs or applications. The API supports a wide range of programming languages and platforms, making it a versatile and convenient tool for a wide range of scientific, medical, and research applications. The API is designed to be flexible, allowing developers to specify the data they need and the format they want it in, while also providing a variety of options for filtering, sorting, and transforming data to meet their specific needs. Overall, the Entrez API provides a powerful and flexible tool for accessing NCBI data and integrating it into a wide range of scientific and medical applications.

The final output of the stage is a folder full of FASTA files that we will use to build the tree.

3.3 Data processing

As soon as the FASTA files were gathered, there are four necessary steps to make this FASTA files operable.

The initial procedure involves assigning new, more descriptive labels to each sequence contained within each FASTA file, thereby rendering the information contained within each sequence more readily comprehended.

The second step entails the alignment of each FASTA file, which will be executed utilizing the MAFFT software, which is a software tool for multiple sequence alignment. It can align large numbers of sequences efficiently and accurately, making it a popular choice in the field of molecular biology and genetics.

The third operation involves the meticulous concatenation of all FASTA files, culminating in the creation of a comprehensive and unified file. To make this possible we used only python dictionaries. This procedure paves the way for conducting a maximum likelihood tree analysis, yet to undertake a Bayesian tree analysis, a final step is require.

Finally, the final step involves tranforming the FASTA file into the NEXUS file format, with the aid of the SeqMagick tool, to optimize the data for further analysis, culminating in the creation of a Bayesian phylogenetic tree. SeqMagick is a powerful and flexible software package that allows for efficient manipulation and conversion of multiple sequence alignment (MSA) files in various formats.

3.4 Tree Inference

The end result of this phase will be the production of two distinct phylogenetic trees - one generated through the Maximum Likelihood method and the other, a Bayesian tree.

To construct the maximum likelihood tree, it was imperative to utilize a FASTA file and the software of choice was RaxML. RaxML is a popular open-source software that uses maximum likelihood algorithms to construct phylogenetic trees from molecular sequence data. The implementation of RaxML within this project allowed for the efficient and accurate creation of the maximum likelihood tree.

The final step in this phylogenetic tree building procedure will entail utilizing the NEXUS file and the MrBayes software to construct a Bayesian tree. A brief introduction to MrBayes, a widely-utilized software for Bayesian inference of phylogenetic trees, must also be provided to contextualize its application in this process.

4 Implementation

The implementation phase is a crucial step in the development process where the methodology and techniques outlined in the previous stages are put into action. This phase involves executing the steps and methods that were designed and tested in the planning phase, resulting in the creation of a functional product or solution. The implementation phase, much like its predecessor, the methodology phase, will encompass four distinct stages of execution.

4.1 Data acquisition

To build the trees, the user was required to provide four arguments to the program, each of which helped to identify the specific combination of folders that were needed for the analysis. These arguments were crucial to ensuring that the correct data was used for each tree, and included the following:

Scientific name of the species - This argument provided the scientific name of the species that was being analyzed, allowing the program to identify the relevant data and folders associated with that species.

Taxonomy hierarchy - The taxonomy hierarchy argument was used to specify the hierarchical classification of the species being analyzed, allowing the program to determine the correct data and folders to use in the analysis.

Proximity - The proximity argument indicated the proximity between the sequences being analyzed, it would indicate how closely related two organisms are, the higher the percentage, higher the relationship between organisms.

Similarity - The similarity argument indicated the level of similarity between the sequences being analyzed, is a measure of how alike two or more sequences or organisms are, based on their genetic or physical characteristics. The higher the similarity value, the more similar these two organisms are.

To obtain the specific taxonomy rank, the user was required to provide two key inputs - the name of the species and the desired taxonomy hierarchy. The taxonomy rank refers to the hierarchical classification of organisms within a taxonomic system, and is used to provide insight into the evolutionary relationships between different species.

For example, if the user entered "homo sapiens" as the name of the species and "order" as the desired taxonomy hierarchy, the program would conduct a search for all primates. By restricting the search based on proximity and similarity, the program was able to identify the specific FASTA folder that was needed for the analysis.

To perform this search, the ENTREZ API was used, which provides direct access to the NCBI database. This database contains a wealth of information on the taxonomy and evolution of different species, and is an indispensable tool for researchers in fields such as biology, genetics, and ecology. By utilizing the ENTREZ API, the user was able to streamline the process of identifying the correct data and folders needed for the analysis, saving time and effort and allowing them to focus on the more complex aspects of their research.

4.2 Data Processing

4.2.1 Fasta Handling

After getting the FASTA folder, it was necessary to change the names of each sequence in each FASTA file inside the folder. The list of names was obtained from the Filtred-ScientificNames_list.txt (an output from the data acquisition phase) and using the SeqIO module, a new folder containing FASTA files was generated, with each file named according to the corresponding sequence. An example of a sequence name, which initially appeared as "KC836121.1 Montifringilla ruficollis mitochondrion, complete genome," was transformed to "Montifringilla ruficollis" after processing with the SeqIO module.

4.2.2 Fasta Alignment

The alignment of the FASTA files was accomplished by converting the folder containing the FASTA files into a list, and then using a for loop to iterate through the list and align each FASTA file with the MAFFT command. The aligned FASTA files were stored in a separate folder for easy access and organization.

4.2.3 Fasta Concatenation

The process of concatenating the FASTA files involved reading each file in the folder of aligned FASTA files, line by line. For each file, the names of the sequences were extracted and used as the keys in a dictionary. The sequences themselves were assigned as the values for each key, providing a clear and structured representation of the data.

By using a dictionary to store the sequences, it was possible to easily access and manipulate the data, making further analysis and processing more efficient. The concatenated FASTA file provided a comprehensive and organized representation of all the sequences, ready for further analysis and interpretation. The concatenated FASTA will also be used to build both trees.

4.3 Tree Inference

4.3.1 Maximum Likelihood Tree

Prior to constructing a maximum likelihood tree, it is crucial to determine the most appropriate evolutionary model of DNA or protein sequence evolution. This can be achieved through the use of software tools such as modeltest-ng, which performs model selection and averaging in phylogenetic analysis. modeltest-ng compares multiple models of molecular evolution and calculates the likelihood of each model, based on the Akaike information criterion (AIC) or Bayesian information criterion (BIC). In this particular case, it was necessary to determine if the selected evolutionary model included the "+I" derivation. After knowing if the model included the "+I" derivation, The time had come to construct the tree using the RaxML command line tool. This command was designed to automatically execute a specified number of times using the "-# autoFC" option, with the goal of generating a newick format file. This file, which serves as the output of the RaxML analysis, will then be utilized to visualize the tree and provide valuable insights into the evolutionary relationships between the sequences in the dataset. By executing the RaxML command in this manner, the user was able to perform a robust and efficient analysis to gain a better understanding of the evolutionary history of the sequences.

4.3.2 Bayesian Tree

The construction of the Bayesian tree posed a slightly different challenge than other types of phylogenetic trees. To begin the process, it was necessary to convert the concatenated FASTA file into a NEXUS file, which is a format that MrBayes, a popular Bayesian phylogenetic analysis software, can work with. This conversion was accomplished using SeqMagick, a powerful Python library designed specifically for the manipulation of biological sequences and sequence files.

Once the necessary inputs had been prepared, the final step was to incorporate the MrBayes command lines into the NEXUS file. This step was crucial, as it ensured that all necessary commands would be executed automatically when MrBayes was run. This streamlined the analysis process and reduced the burden of manual intervention, freeing the user to focus on the interpretation and analysis of the results.

By combining the powerful capabilities of SeqMagick and MrBayes, the user was able to efficiently and effectively build a Bayesian tree, gaining valuable insights into the evolutionary relationships between the sequences in the dataset. This type of analysis is critical for understanding the underlying mechanisms driving evolutionary change, and is a critical tool for researchers in many fields, including biology, genetics, and ecology.

4.4 Test Suite

To ensure the accuracy of our results, a comprehensive test suite was implemented throughout the phylogenetic tree building process. The test suite consisted of a series of validation
checks for each step, starting the data acquisition with EntrezAPI, to the final phylogenetic
tree output using MrBayes. The validation checks included verifying the integrity of input
data, examining the alignment quality, and evaluating the consistency of tree topology. The
results of the tests were recorded and analyzed to ensure that the program meets the required standards and produces accurate phylogenetic trees. The test were implement using
pytest. Pytest is a popular testing framework for Python programming language that is
used to write and run tests for applications and libraries. It allows developers to write tests
in a simple, concise and readable manner. Pytest provides a rich set of tools for writing
tests, including fixtures for setup and teardown, assertion introspection and plugins for
extended functionality.

5 Results

- Colocar os mambinhos dos graficos e exemplis de fastas, alignments, concatenate, etc
- Summary of the results obtained from testing the software, including any performance metrics and examples of the generated phylogenetic trees.
- Observations or insights gained from the results, and how they compare to expected or previous results.
- It should provide any visualizations or plots that help to interpret the results and explain any patterns or trends found in the data.

6 Conclusion

References

- [1] Risso D, Schwartz K, Sherlock G, Dudoit S (2011). GC-Content Normalization for RNA-Seq Data. *BMC Bioinformatics*, 12(1), 480
- [2] asdfasdfasdf, 12(1), 480