- <sup>1</sup> Physcraper: a python package for continual update of evolutionary
- estimates using the Open Tree of Life
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## $_{ ilde{0}}$ 1 Abstract

- 21 1. Phylogenies are a key part of research in all areas of biology. Tools that automatize some parts of the
  22 process of phylogenetic reconstruction (mainly character matrix construction) have been developed for
  23 the advantage of both specialists in the field of phylogenetics and nonspecialists. However, interpretation
  24 of results, comparison with previously available phylogenetic hypotheses, and choosing of one phylogeny
  25 for downstream analyses and discussion still impose difficulties to one that is not a specialist either on
  26 phylogenetic methods or on a particular group of study.
- 2. Physcraper is an open-source, command-line Python program that automatizes the update of published phylogenies by making use of public DNA sequence data and taxonomic information, providing a framework for comparison of published phylogenies with their updated versions.
- 3. Physcraper can be used by the nonspecialist, as a tool to generate phylogenetic hypothesis based on
  already available expert phylogenetic knowledge. Phylogeneticists and group specialists will find it useful
  as a tool to facilitate comparison of alternative phylogenetic hypotheses (topologies). Is physcraper
  intended for the nonspecialist?? We have two types of nonspecialists: the ones that do
  not know about phylogenetic methods and the ones that might know about phylogenetic
  methods but do not know much about a certain biological group.
- 4. Physcraper implements node by node/topology comparison of the the original and the updated trees using the conflict API of OToL, and summarizes differences.
- 5. We hope the physcraper workflow demonstrates the benefits of opening results in phylogenetics and encourages researchers to strive for better data sharing practices.
- 6. Physcraper can be used with any OS. Detailed instructions for installation and use are available at https://github.com/McTavishLab/physcraper.
- 42 **Keywords**: cross-connectivity, gene tree, open science, open tree of life, phylogeny, public database, python,
- reproducibility, taxonomy, updated alignment

## 4 2 Introduction

- 45 From molecular data to alignments and phylogenies, public biological data repositories such as the GenBank
- database (Benson et al. 2000; Wheeler et al. 2000), the TreeBASE repository (Piel et al. 2009) and the Open
- <sup>47</sup> Tree of Life (OToL) data deposition tool, the phylesystem (McTavish et al. 2015), are still accumulating data.
- 48 More than a decade ago, the GenBank database release number 159 (April 15, 2007) had 72 million DNA
- 49 sequences, and was estimated to have the potential to resolve phylogenetic relationships of about 98.05%
- of taxa included in the NCBI taxonomy (Sanderson et al. 2008) which represents about 10% of described
- 51 diversity (Federhen 2003; Scott 2011), taking a conservative estimate of extant diversity. In comparison, the
- $_{52}$  current GenBank release number 238 (June 15, 2020) has tripled in size, hosting data for more than 217
- million DNA sequences https://ftp.ncbi.nih.gov/genbank/gbrel.txt.
- Many useful tools have been developed in the past decades in an effort to make sense of the large amount of
- 55 DNA data in public and private databases. Most of these tools were motivated by the genomics revolution,
- to identify clusters of homologs for genomic assembly. Notably, this homolog DNA clusters can be used
- 57 as homology hypotheses to reconstruct phylogenetic relationships. Tools that automatize the assembly of
- box homology hypotheses from the GenBAnk database for phylogenetics such as PHYLOTA (Sanderson et al.
- 59 2008), PHLAWD (Smith et al. 2009), and SUPERSMART (Antonelli et al. 2017) have been widely used to
- 60 reconstruct the evolutionary relationships among different organisms.
- 61 This has been an important contribution to the field of phylogenetics, especially be they represent a possible
- 62 way for reproducibility in phylogenetics. Indeed, most published phylogenies have been infered from a
- 63 homology hypothesis that has been curated "by hand" (Morrison 2009).
- on one hand, there are concerns about the use of automated tools. Concerns I think people have about these
- tools: Errors in identification of sequences Little control along the process Too much of a black box?
- 66 Most of these phylogenies are being constructed by people learning about the methods (aka students), so
- 67 they want to know what is going on at each step and end up doing it manually. It has also been suggested
- that manual curation of genetic datasets always produces better results in phylogenetic analyses.

- <sup>69</sup> Fortunately, public manually curated homology hypotheses have the potential to be enriched and updated by
- incorporating newly released data from public molecular databases as an alternative to automatic identification
- and clustering of DNA homologs for assembly of homology hypotheses to reconstruct phylogenetic relationships.
- 72 There are THIS MANY curated alignments publicly available on TreeBASE
- 73 This suggests a lot of potential for update of phylogenetic relationships in different regions of the tree of life.
- OToL has integrated the phylogenies fro TreeBASe, and linked to the alignment repo but not the exact
- <sup>75</sup> alignment that generated the phylogeny.
- Linkage of original alignment with corresponding phylogeny has t be done by a human curator.
- 77 We present a way to automatize and standardize the comparison of phylogenetic hypotheses and to allow
- reproducibility of this last step of the research process.
- 79 A key aspect of the standard phylogenetic workflow is comparison with already existing phylogenetic hypotheses
- 80 and with phylogenies that are considered "best" by experts not only in phylogenetics, but also experts on the
- 81 focal group of study.
- 82 The pipelines are so powerful and they will give you an answer, but there is no way to assess if it is better
- than previous answers, it just assumes it is better because it used more data.
- The goal of Physcraper is to build upon previous phylogenetic knowledge, allowing a direct comparison
- between existing phylogenies and phylogenies that are constructed using new genetic data retrieved from a
- public nucleotide database (i.e., GenBank (Benson et al. 2000; Wheeler et al. 2000)).
- 87 To achieve this, Physcraper uses the Open Tree of Life phylesystem and connects it to the TreeBase database,
- to (1) get the original DNA data set matrices (alignments) that produced a phylogeny that was published
- and then made available in the OToL database, (2) use this DNA alignments as a starting point to get new
- genetic data belonging to the focal group of study, to (3) finally update the phylogenetic relationships in the
- group.

- 92 A less automated workflow is one in which the alignments that generated the published phylogeny are stored
- 93 in other public database (such as DRYAD) or elsewhere (the users computer), and are provided by the users.
- 94 Physcraper implements node by node comparison of the the original and the updated trees, using the conflict
- 95 API of OToL.

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# <sup>96</sup> 3 How does Physcraper work?

#### 3.1 The input: a study tree and an alignment

- The study tree is a published phylogenetic tree stored in the OToL database, phylogystem (McTavish et al. 2015). The main reason for this is that trees in phylogystem have a set of user defined characteristics that are essential for automatizing the phylogeny update process. The most relevant of these being the definition of ingroup and outgroup. Outgroup and ingroup taxa in the original tree are identified and tagged. This allows to automatically set the root for the updated tree on the next steps of the pipeline.

  A user can choose from the 'r rotl::tol\_about()\$num\_source\_trees' published trees supporting the resolved node of the synthetic tree in the OToL website (<>). If the tree you are interested in updating
  - The alignment should be a gene alignment that was used to generate the tree. The original alignments are usually stored in a public repository such as TreeBase (Piel et al. 2009; Vos et al. 2012), DRYAD (http://datadryad.org/), or the journal were the tree was originally published. If the alignment is stored in TreeBase, physcraper can download it directly, either from the TreeBase website (https://treebase.org/) or through the TreeBase GitHub repository (SuperTreeBase; https://github.com/TreeBase/supertreebase). If the alignment is on another repository, or provided personally by the owner, a copy of it has to be downloaded by the user, and it's local path has to be provided as an argument.

is not in there, you can upload it via OToL's curator tool (<a href="https://tree.opentreeoflife.org/curator">https://tree.opentreeoflife.org/curator</a>).

- A taxon name matching step is performed to verify that all taxon names on the tips of the tree are in the DNA character matrix and vice versa.
- A ".csv" file with the summary of taxon name matching is produced for the user.

- Unmatched taxon names are dropped from both the tree and alignment. Technically, just one matching

  name is needed to perform the searches. Please, see next section.
- A ".tre" file and a ".fas" file containing only the matched taxa are generated and saved in the inputs

  folder to be used in the following steps.

## 3.2 DNA sequence search and cleaning

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- The next step is to identify the search taxon within the reference taxonomy. The search taxon will 122 be used to constraint the DNA sequence search on the nucleotide database within that taxonomic 123 group. Because we are using the NCBI nucleotide database, by default the reference taxonomy is the NCBI taxonomy. The search taxon can be provided by the user. If none is provided, then 125 the search taxon is identified as the Most Recent Common Ancestor (MRCA) of the matched taxa 126 belonging to the ingroup in the tree, that is also a named clade in the reference taxonomy. This 127 is known as the Most Recent Common Ancestral Taxon (MRCAT; also referred in the literature 128 as the Least Inclusive Common Ancestral Taxon - LICA). The MRCAT can be different from the 129 phylogenetic MRCA when the latter is an unnamed clade in the reference taxonomy. To automatically 130 identify the MRCAT of a group of taxon names, we make use of the OToL taxonomy tool (https: 131 //github.com/OpenTreeOfLife/germinator/wiki/Taxonomy-API-v3#mrca). 132
  - Users can provide a search taxon that is either a more or a less inclusive clade relative to the ingroup of the original phylogeny. If the search taxon is more inclusive, the sequence search will be performed outside the MRCAT of the matched taxa, e.g., including all taxa within the family or the order that the ingroup belongs to. If the search taxon is a less inclusive clade, the users can focus on enriching a particular clade/region within the ingroup of the phylogeny.
- The Basic Local Alignment Search Tool, BLAST [Altschul et al. (1990); altschul1997gapped] is used
  to identify similarity between DNA sequences within the search taxon in a nucleotide database, and
  the accepted sequences on the alignment. The blastn function from the BLAST command line tools
  (Camacho et al. 2009) is used for local-database searches. A modified biopython blast function is used

- for web-based searches.
- The DNA sequence similarity search can be done on a local database that is easily setup by the user.

  In this case, the blastn function is used to performs the similarity search (Camacho *et al.* 2009).
- The search can also be performed remotely, on the NCBI database. In this case, the bioPython BLAST function was modified to accepts is used to perform the similarity search.
- A pairwise alignment-against-all BLAST search is performed. This means that each sequence in the
  alignment is BLASTed against DNA sequences in a nucleotide database constrained to the search taxon.
  Results from each one of these BLAST runs are recorded, and matched sequences are saved along with
  their corresponding identification numbers (accession numbers in the case of the GenBank database).
  This information will be used later to store the whole sequences in a dedicated library within the
  physcraper folder, allowing for secondary analyses to run significantly faster.
- Matched sequences below an e-value, percentage similarity, and outside a minimum and maximum length
   threshold are discarded. REPORT THE DEFAULT VALUES AND DESCRIBE WHAT
   THEY MEAN This filtering leaves out genomic sequences. All acepted sequences are asigned an
   internal identifier, and are further filtered.
- Because the original alignments usually lack database id numbers, a filtering step is needed. Accepted
  sequences that belong to the same taxon of the query sequence, and that are either identical or shorter
  than the original sequence are discarded. Only longer sequences belonging to the same taxon as the
  original sequence will be considered further for analysis.
- Among the remaining filtered sequences, there are usually several exemplars per taxon. Although it
  can be useful to keep some of them to, for example, investigate monophyly within species, there can be
  hundreds of exemplar sequences per taxon for some markers. To control the number of sequences per
  taxon in downstream analyses, 5 sequences per taxon are chosen at random. This number is set by
  default but can be modified by the user.

- Reverse complement sequences are identified and translated.
- Users can choose to perform a more "cycles" of sequence similarity search, by blasting the newly found
  sequences. This can be done iteratively, but by default only sequences in the alignment are blasted. Is
  there an argument to control the number of cycles of blast searches with new sequences?
- Accepted sequences are downloaded in full, and stored as a local database in a directory that is globally accesible (physcraper/taxonomy), so they are accesible for further runs.
- A fasta file containing all filtered and processed sequences resulting from the BLAST search is generated for the user.

#### 174 3.3 DNA sequence alignment

- The software MUSCLE (Edgar 2004) is implemented to perform alignments.
- First, all new sequences are aligned using default MUSCLE options.
- Then, a MUSCLE profile alignment is performed, in which the original alignment is used as a template
  to align new sequences. This ensures that the final alignment follows the homology criteria established
  by the original alignment.
- The final alignment is not further processed automatically. We encourage users to check it either by eye and perform manual refinement or using any of the many tools for alignment processing, to eliminate columns with no information.

#### 3.4 Tree reconstruction and comparison

- A gene tree is reconstructed for each alignment provided, using a Maximum Likelihood approach implemented with the software RAxML (Stamatakis 2014) with 100 classic rapid bootstrap (Felsenstein 1985) replicates by default. The number of bootsrap replicates can be modified by the user. Other type of bootstrap that I think is not yet incorporated into physcraper is the Transfer Bootstrap Expectation (TBE) recently proposed in Lemoine et al. (2018).
- The original tree is used as starting tree for the ML searches. It can also be set as a full topological

- constraint or not used at all, depending on the goals of the user.
- Bootstrap results are summarized with Dendropy ADD CITATION
- The final result is an updated phylogenetic hypothesis for each of the genes provided in the alignment.
- Tips on all trees generated by physcraper are defined by a taxon name space, allowing to perform comparisons and conflict analyses.
- Robinson Foulds weighted and unweighted metrics ARE CALCULATED WITH DENDROPY TOO.
- Describe what a conflict analysis is: Node by node comparison of the resulting clades compared to

  CITE REDELINGS AND HOLDER (??? and holder)
- For the conflict analysis to be meaningful, the root of the tree ineeds to be accurately defined.
- A SUGGESTED DEFAULT ROOTING BASED ON THE OPEN TREE TAXONOMY is implemented

  for now. DESCRIBE HOW IT WORKS. SAY THAT IT IS A PROBLEM. Automatic rooting is not

  that smart yet. The best way right now is for users to define outgroups so trees are better rooted.
- Currently, the root is determined by finding the parent node of the sequences that do not belong to the ingroup/ search taxon. This ensures a correct rooting of the tree even when the search taxon is more inclusive than the ingroup.
- Conflict information can only be generated in the context of the whole Open Tree of Life. Otherwise, it is not really possible to get conflict data. One way to compare two independent phylogenetic trees is to compare them both to the synthetic OToL and then measure how well they do against each other

# $_{\scriptscriptstyle{209}}$ 4 Examples

To exemplify the utility of physcraper we will address two use-case scenarios. One in which the user is interested in a particular group. Another one in which the user is interested in a particular phylogeny. A tutrial as well as illustrated examples of steps needed to perform a physcraper analyses are available elsewhere.

#### $_{213}$ 4.1 The hollies

A student is interested in the genus *Ilex*, the only extant clade within the family Aquifoliaceae, order Aquifoliales of flowering plants. The genus encompasses between 400-600 living species. A review of literature reveals that there are three phylogenetic trees showing relationships within the hollies published. The first one has been made available in TreeBASE as well as in the OToL phylesystem and is part of the synthetic tree. 217 It samples 48 species. The second tree has not been made available anywhere, not even in the supplementary 218 data of the original publication. The most recent one has been made available in the OToL Phylesystem and in the DRYAD repository. It is the best sampled yet, with 200 species. However, it has not been added to 220 the syntehtic tree yet. This makes it a perfect case to test the basic functionalities of physcraper: we know 221 that the sequences of the most recently published tree have been made available on the GenBank database. 222 Hence, we expect that updating the oldest tree will produce something very similar to the newest tree. 223

#### 224 4.2 The Malvaceae

A postdoc started working with a new reserach group. They are interested in solving relationships among lineages of the Malvaceae, a family of flowering plants with almost 6 000 known species. A review of the literature shows them that there are many phylogenetic trees encompassing some of the linegaes in the group. However, the head of the group wants to use a particular marker they beilieve to be the best one to be able to solve the relationships in the group. They have been working in the alignment for long and they want to incorporate new data into the hypothesis of homology they have been curating and that they trust.

# Original tree Updated tree

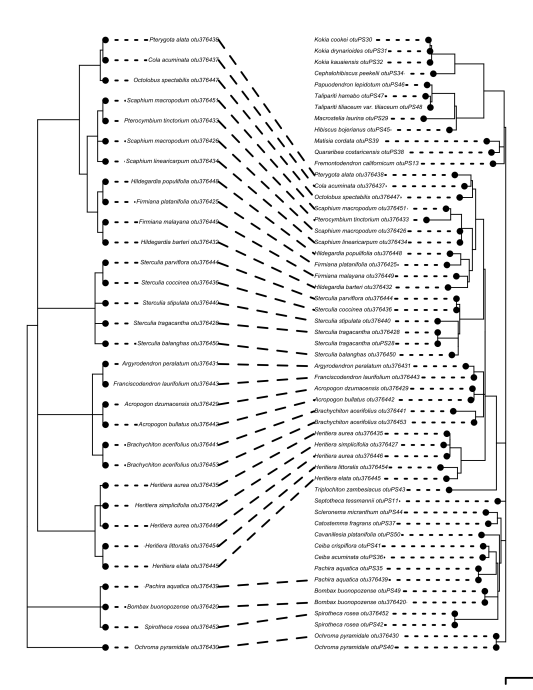


Figure 1: Comparison of original tree and update tree of the Malvaceae.

## 5 Discussion

- Data repositories hold more information than meets the eye. Besides the actual data, they have other types of information that can be used for the advantage of science.
- Usually, initial ideas about the data are changed by analyses. We expect that this new ideas on the data can
  be registered on data bases, exposing new comers to expert understanding about the data.
- There are many tools that are making use of DNA data repositories in different ways. Most of them focus on
  efficient ways to mine the data getting the most homologs. Some focus on accurate ways of mining the data
   getting real and clean homologs. Others focus on refinement of the alignment. Most focus on generating full
  trees de novo, mainly for regions of the Tree of Life that have no phylogenetic assessment yet in published
  studies, but also for regions that have been already studied and that have phylogenetic data already.
- All these tools are great efforts for advancing towards reproducibility in phylogenetics, a field that has been largely recognised as somewhat artisanal. We propose adding focus to other sources of information available from data repositories. Taking advantage of public DNA data bases have been the main focus. However, phylogenetic knowledge is also accumulating fast in public and open repositories. In this way, the physcraper pipeline can be complemented with other tools that have been developed for other purposes.
- We emphasize that physcraper takes advantage of the knowledge and intuition of the expert community to
  build upon phylogenetic knowledge, using not only data accumulated in DNA repositories, but phylogenetic
  knowledge accumulated in tree repositories. This might help generate new phylogenetic data. But physcraper
  does not seek to generate full phylogenies de novo.
- Describe again statistics to compare phylogenies provided by physcraper via OpenTreeOfLife. Mention statistics provided by other tools: PhyloExplorer (Ranwez *et al.* 2009). Compare and discuss.
- How is physcraper already useful: to mine targeted sequences, in this way it is similar to baited analyses
  from PHLAWD and pyPHLAWD. Phylota does not do baited analyses, I think, only clustered analyses. Finding

How can it be used for the advantage of the field: - rapid phylogenetic placing of newly discovered species, as mentioned in Webb et al. (2010) - obtain trees for ecophylogenetic studies, as mentioned in Helmus & Ives 256 (2012) - one day could be used to sistematize nucleotide databases, such as Genbank (Benson et al. 2000; 257 Wheeler et al. 2000), as mentioned in San Mauro & Agorreta (2010), i.e., curate ncbi taxonomic assignations. 258 allows to generate custom species trees for downstream analyses, as mentioned in Stoltzfus et al. (2013) Things that physcraper does not do: - analyse the whole GenBank database (Benson et al. 2000; Wheeler 260 et al. 2000) to find homolog regions suitable to reconstruct phylogenies, as mentioned in Antonelli et al. 261 (2017). There are already some very good tools that do that. - provide basic statistics on data availability to 262 assemble molecular datasets, as mentioned by Ranwez et al. (2009). Phyloexplorer does this? - it is not a 263 tree repo, as phylota is, mentioned in Deepak et al. (2014) 264

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## <sup>269</sup> 7 Authors' Contributions

# 270 8 Data Avilability

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