- Physcraper: a python package for continual update of evolutionary
- estimates using the Open Tree of Life
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Abstract

- 1. Phylogenies are a key part of research in many areas of biology. Tools that automate some parts of
 the process of phylogenetic reconstruction, mainly molecular character matrix assembly, have been
 developed for the advantage of both specialists in the field of phylogenetics and nonspecialists. However,
 interpretation of results, comparison with previously available phylogenetic hypotheses, and choice
 of one phylogeny for downstream analyses and discussion still impose difficulties to one that is not a
 specialist either on phylogenetic methods or on a particular group of study.
- 2. Physcraper is a command-line Python program that automates the update of published phylogenies by adding public DNA sequences to underlying alignments of previously published phylogenies. It also provides a framework for straightforward comparison of published phylogenies with their updated versions, by leveraging upon tools from the Open Tree of Life project to link taxonomic information across databases.
- 33. Physcraper can be used by the nonspecialist, as a tool to generate phylogenetic hypotheses based on
 publicly available expert phylogenetic knowledge. Phylogeneticists and taxonomic group specialists
 will find it useful as a tool to facilitate molecular dataset gathering and comparison of alternative
 phylogenetic hypotheses (topologies).
- 4. The Physcraper workflow demonstrates the benefits of doing open science for phylogenetics, encouraging
 more researchers to strive for better sharing practices. Physcraper can be used with any OS and is
 released under an open-source license. Detailed instructions for installation and use are available at
 https://physcraper.readthedocs.
- 41 **Keywords**: gene tree, interoperability, open science, open tree of life, phylogeny, public database, python,
- reproducibility, taxonomy, updated alignment

1 Introduction

Phylogenies capture the shared history of living and extinct organisms and provide key evolutionary context for all our biological observations. Public biological databases constitute an amazing resource for evolutionary studies, but a large portion of molecular data publicly available has never been incorporated into any phylogenetic estimate. Extending existing phylogenies with new DNA sequence data, geographical location, and other metadata in a reproducible and continuous manner is possible by automating connections between biological databases. Here, we introduce Physcraper, a tool to build upon molecular data that taxon specialists have assessed and deemed appropriate for a specific phylogenetic scope, by using sequence data from public biological databases to update a starting tree and single locus alignments. Automatic integration of data in a phylogenetic context is challenged by the prevalence of taxonomic idiosyncrasies across biological databases. Taxonomic name standardization represents a step toward tackling this issue, but requires a unified system. The Open Tree of Life project (OpenTree) has constructed a comprehensive tree of life by synthesizing published phylogenies along with taxonomic data. OpenTree's "synthetic" tree comprises 2.3 million tips, of which around 90,000 are supported by phylogenetic estimates the remaining 1.4 million taxa are placed in the tree based on taxonomy. To achieve this, OpenTree unifies taxonomy from various databases (Rees & Cranston 2017), including the USA National Center for Biodiversity Information (NCBI) molecular database GenBank (Benson et al. 2000), the Global Biodiversity Information Facility (GBIF), the World Register of Marine Species (WoRMS), and other resources. The OpenTree unified taxonomy represents a key resource for connecting data from virtually any biological database to phylogenetic data that has been standardized to OpenTree's unified taxonomy. Another challenge to incorporating molecular data from public databases to update phylogenetic knowledge is assembling high-quality homology hypotheses. Species tree reconstructions from multiple single locus data sets taking into account the multispecies coalescent model are considered the gold standard for inferring species relationships (Song et al. 2012). Genomics has, and will continue to, revolutionize phylogenetic inference. Yet, different research questions call for different genomic sequencing approaches (e.g., whole genomes vs microsatellites), which produce largely non-overlapping genomic data sets across taxa, creating challenges in wide scale phylogenetic reconstruction. While phylogenomics ameliorates the problem of non-overlapping genomic data sets by focusing on targeted capture of informative characters from independent and single-copy genetic markers (Andermann *et al.* 2020), decades of single locus sequencing have generated massive amounts of homologous DNA data sets that can be used for phylogenetic reconstruction at many scales. Even in phylogenomic reconstructions, thoughtfully curated markers and alignments can improve phylogenetic reconstructions (Fragoso-Martínez *et al.* 2017).

More than a decade ago, GenBank release number 159 (April 15, 2007) already hosted 72 million DNA sequences. These sequences were gauged to have the potential to resolve phylogenetic relationships of most (98.05%) of the almost 241, 000 distinct taxa in the NCBI taxonomy at the time (Sanderson et al. 2008). Assembling a DNA alignment from such a massive database can be done "by hand", but it is a largely time consuming and mostly non-reproducible approach. Computational pipelines that make DNA sequence search faster and more efficient, as well as more reproducible, have been applied to study evolutionary relationships among a variety of organisms (e.g., Smith et al. 2009; Izquierdo-Carrasco et al. 2014; Antonelli et al. 2017).

A way to incorporate the benefits from massive amounts of newly released molecular data and fine-grained curation from human experts, is to rely on manually curated homology hypotheses as "jump-start" alignments (Morrison 2006). The TreeBASE database (Piel et al. 2009) hosts about 8, 200 alignments, providing information on evolutionary relationships of around 100, 000 distinct taxa (see TreeBASE's website about), representing a public source of valuable expert knowledge. Linking published alignments with molecular data that has not yet been included in any public phylogenetic estimate, has the potential to accelerate the enrichment and updating of phylogenetic relationships in many regions of the tree of life.

Physcraper relies on programmatic access protocols (API's) to automatically link molecular data from GenBank to alignments from TreeBASE and phylogenies from OpenTree's Phylogenetic, to continually update and enrich phylogenetic knowledge based on expertly-curated homology hypotheses. Physcraper also provides new types of access to various OpenTree tools for comparison of existing phylogenetic hypotheses with newly generated ones. Physcraper is coded as a Python pipeline that focuses on data interoperability, by using the standardized taxonomy as a way to link taxon data from different databases. This integration also allows

- users to rapidly place new data from a diverse range of biological databases in an evolutionary context,
- opening the possibility for a variety of comparative downstream analyses.

₉₇ 2 The Physcraper framework

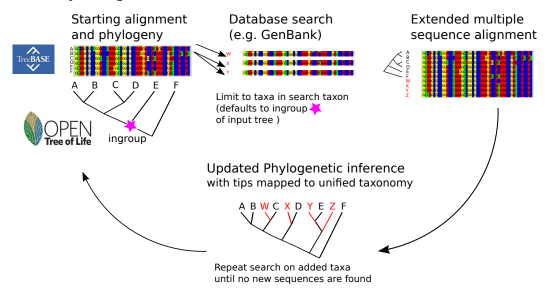


Figure 1: The Physcraper framework consists of 4 steps (see text). The software is fully described on its documentation website at physcraper.readthedocs.io, along with installation instructions, function usage descriptions, examples and tutorials.

- The general Physcraper framework is depicted in Figure 1. It consists of 4 steps: 1) identifying and processing
- ⁹⁹ a phylogenetic tree to update its underlying alignment; 2) performing a constrained BLAST search of
- 100 DNA sequences in the original alignment on the GenBank database, and filtering of new sequences; 3)
- profile-aligning filtered new sequences to the original alignment; 4) performing a phylogenetic analysis and
- 102 comparing the updated tree to previous phylogenetic estimates within the focus group.

103 2.1 The inputs: a tree and an alignment

- Taxon names in the input tree must be standardized or "mapped" to the unified OpenTree taxonomy (Rees
- 8 Cranston 2017) using OpenTree's bulk Taxonomic Name Resolution Service TNRS tool. Users can upload
- their own tree, or choose from among the 2, 950 mapped trees stored in OpenTree's Phylesystem that also
- have alignments avilable on TreeBASE (Piel et al. 2009).

The input alignment should be a single locus alignment that was used in part or in whole, to generate the input tree. If the alignment associated to the input tree is stred in TreeBASE, Physcraper retrieves it automatically.

Alignments stored in any other repository, or constituting personal data have to be downloaded by the user.

Physcraper processes the input tree to match taxa found in the alignment, and verifies that all taxon names on the tips of the tree are in the DNA character matrix and vice versa. Technically, just one taxon name and its corresponding sequence in the alignment are needed to continue the algorithm.

114 2.2 DNA sequence search and filtering

The DNA sequence search is performed with the Basic Local Alignment Search Tool, BLAST (Altschul et al. 1990), either on the GenBank remote database or in a GenBank local database set up by the user. It is constrained to a taxonomic group in the NCBI taxonomy, defined as the "search taxon". Users can arbitrarily define a search taxon that is either a more or a less inclusive clade relative to the ingroup of the input tree. Otherwise, the search taxon is automatically identified using the OpenTree API (Rees & Cranston 2017), as the Most Recent Common Ancestral Taxon (MRCAT) of the ingroup taxa in the input tree, i.e., the MRCA that is also a named clade in the NCBI taxonomy (Fig. 1). The MRCAT can be different from the phylogenetic MRCA when the latter is an unnamed clade in the synthetic tree.

BLAST command line tools blastn function (Camacho et al. 2009) is used for local database sequence
searches. For remote database searches, we modified the BioPython (Cock et al. 2009) BLAST function
from the NCBIWWW module to accept an alternative BLAST address (URL). This is useful when a user
lacks access to the computer capacity needed to setup a local database, but has access to an institutional
server. Each sequence in the alignment is BLASTed once against all DNA sequences in GenBank within the
search taxon. GenBank sequences with match scores smaller than the e-value cutoff (default to 0.00001) are
downloaded into a local library.

Downloaded sequences are not further considered in the analysis if they fall outside a min and max length threshold, defined as the proportion of the average length without gaps of all sequences in the input alignment (default values of 80% and 120%, respectively); or if they are either identical to or shorter than an existing

- sequence in the input alignment and they represent the same taxon in the OTT taxonomy or the NCBI taxonomy. If there are several sequences per taxon, an arbitrary number of 5 (can be modified by the user) sequences per taxon are chosen at random.
- Reverse, complement, and reverse-complement sequences are identified and translated using BioPython internal functions (Cock *et al.* 2009). Iterative cycles of BLAST searches can be performed, by blasting the new sequences until no new ones are found. By default only one BLAST search cycle is performed in which only sequences in the input alignment are blasted.

140 2.3 New DNA sequence alignment

- By default, Physcraper uses MUSCLE (Edgar 2004) to perform DNA sequence alignments in a two step process. First, all new sequences are aligned using the default MUSCLE options.
- Second, a MUSCLE profile alignment is performed, in which the original alignment is used as a template to align the new sequences. This ensures that the final alignment follows the homology criteria established by the original alignment. The final alignment is not further processed by Physcraper. It is recommended that the alignment is checked by the user, by eye followed by manual refinement, or using a tool for automatic alignment processing (e.g., GBlocks; Castresana 2000). Users may also use Physcraper to only gather new GenBank sequences, to then apply their own preferred alignment and phylogenetic inference methods.

149 2.4 Tree reconstruction and comparison

- A Maximum Likelihood (ML) gene tree is reconstructed for each alignment provided, using RAxML (Stamatakis 2014) with default settings (GTRCAT model of molecular evolution and 100 bootstrap replicates with the default algorithm). Only the number of bootsrap replicates can be defined by the user. By default, the original tree is used as a starting tree for the ML searches. Alternatively, the original tree can be set as full topological constraint, or simply be ignored for the ML searches. Bootstrap results are summarized with the SumTrees module of DendroPy (current version 4.4.0; Sukumaran & Holder 2010).
- Physcraper's main result is an updated phylogenetic hypothesis for the search taxon. The updated tree can

be compared with the original tree using an automated conflict analysis which calculates Robinson Foulds weighted and unweighted metrics using Dendropy (Sukumaran & Holder 2010), and performs a node by node 158 comparison between the synthetic OpenTree and the original and updated tree individually, using OpenTree's 159 conflict API (Redelings & Holder 2017). For the conflict analysis to be meaningful, the root of the tree needs 160 to be accurately defined. A default rooting based on OpenTree's taxonomy is implemented for now. It uses 161 the taxon labels for all the tips in the updated tree, pulls an inferred subtree from OpenTree's taxonomy and 162 then applies the same rooting to the inferred updated tree. However, if the updated tree changes expectations 163 from taxonomy, the rooting may no longer be appropriate. Automatic identification of a phylogenetic tree root is a difficult problem that has not been solved yet. The best way right now is for users to define the outgroup directly on the updated tree as part of the conflict analysis, so trees are accurately rooted.

₆₇ 3 Case Study: The hollies

A user is interested in the phylogenetic relationships within the genus *Ilex*. Commonly known as "hollies", the genus encompasses between 400-700 living species, and is the only extant clade within the family Aquifoliaceae, order Aquifoliales of flowering plants.

An online literature review in June 2020 (google scholar search for "ilex phylogeny") reveals that there are several published phylogenetic trees showing relationships within the hollies (Cuénoud et al. 2000; Setoguchi & Watanabe 2000; Selbach-Schnadelbach et al. 2009; Manen et al. 2010), but only two have their data available publicly (Gottlieb et al. 2005; Yao et al. 2020). Gottlieb et al. (2005) made original tree and alignment data available in TreeBASE. The "Gottlieb2005" tree sampling 41 species was added to the OpenTree Phylesystem and its information has been integrated into OpenTree's synthetic tree.

The most recent *Ilex* tree from Yao *et al.* (2020), has been made available in the OpenTree Phylesystem and in the DRYAD repository. The "Yao2020" tree, is the best sampled phylogenetic tree yet available for the hollies, with 175 tips.

80 A tutorial as well as illustrated examples of functions implemented on each step of the analysis are available

in Physcraper's documentation website. Figure 2 shows results from the Physcraper analysis of an alignment 181 of the internal transcribed spacer DNA region (ITS) from Gottlieb et al. (2005). Physcraper ran on a local 182 BLAST database, on a laptop Linux computer for 19hrs 45min to perform BLAST and RAxML analyses, 183 with bootstrap analyses taking an additional 13hrs. The updated Gottlieb2005 tree contains all 41 distinct 184 taxa from the original study plus 231 new tips, contributing phylogenetic data to 84 additional *Ilex* taxa. 185 The best RaxML tree is 99% resolved, with 25% of nodes with bootstrap support < 0.1 and 48% nodes with 186 bootstrap support > 0.75. A large portion of internal branches are negligibly small, with 30 branches < 187 0.00001 substitution rate units, from which only 9 have a bootstrap support > 0.75 (Fig. 2). For comparison, the Yao2020 tree also contains all 41 distinct taxa from the original Gottlieb2005 study, and contributes phylogenetic data to 134 additional *Ilex* taxa, from which 67 are also in the Physcraper updated Gottlieb2005 tree. While Yao et al. (2020) also used ITS as a marker, their data in GenBank is not public yet, so 191 Physcraper was unable to incorporate 68 additional taxa into the analysis. However, Physcraper was able to incorporate 18 taxa that were not in Yao2020. This might be caused by the method they used to download 193 existing ITS Ilex sequences from GenBank, which is not fully explained in the publication, but seems to be a 194 "manual" process.

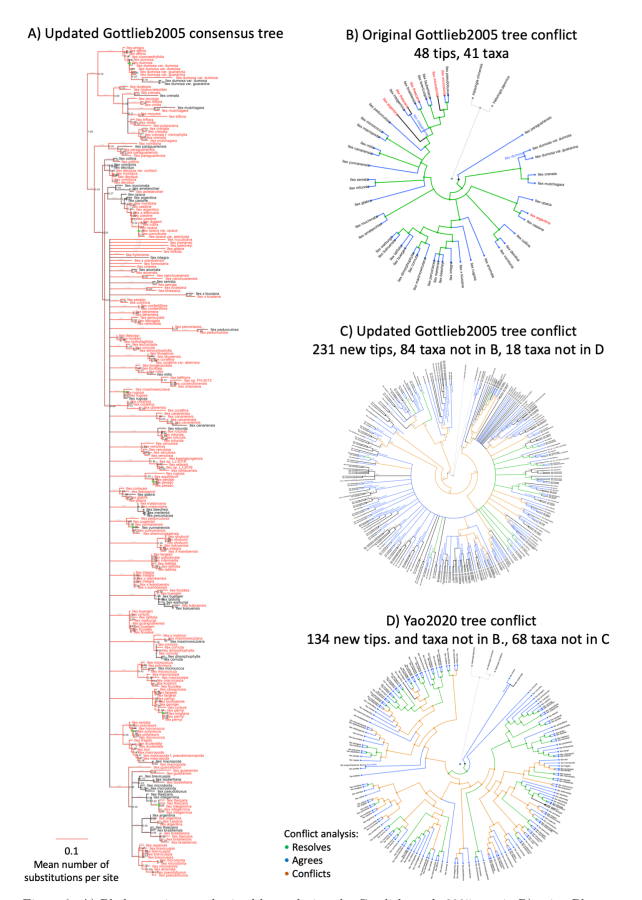


Figure 2: A) Phylogenetic tree obtained by updating the Gottlieb et al. 2005 tree in B) using Physcraper.

Figure 2 caption continued: Tips in original alignment and new tips added with Physcraper are depicted in
black and red, respectively. Physcraper obtained sequences from the GenBank database via local BLAST of
all sequences in the original alignment that generated tree in B), filtered them following criteria specified in
section "DNA sequence search and filtering", aligned them to the original alignment using MUSCLE and
performed a phylogenetic reconstruction using RAxML with 100 bootstraps. B-D show results of the conflict
analysis comparing estimated relationships to taxonomy, performed with OpenTree tools.

₂ 4 Discussion

Data repositories designed to preserve and democratize access to biological data are essential resources for evolutionary estimation. While data keep accumulating, our ability to reanalyze and incorporate new data with available knowledge is being challenged.

While phylogenetic pipelines designed to make evolutionary sense of the vast amount of public molecular 206 data are available (e.g., Phylota (Sanderson et al. 2008), PyPHLAWD (Smith et al. 2009), SUPERSMART 207 (Antonelli et al. 2017)), they focus on generating full trees de novo (i.e., inferring phylogenetic relationships 208 from a newly generated homology hypothesis, as opposed to e.g., supertrees, that are generated by assembling 209 previous phylogenetic estimates). While Physcraper does not generate phylogenies de novo in a traditional 210 sense, it successfully generates new phylogenetic knowledge, revealing the potential of phylogenetic knowledge 211 published in databases to facilitate phylogenetic placement of public molecular data. The PUMPER pipeline 212 (Izquierdo-Carrasco et al. 2014) also uses the concept of updating pre-existing alignments to incorporate public molecular data into phylogenies. Unfortunately, installation of the tool was unsuccesful following instructions from the author, and we were unable to benchmark a comparison.

Physcraper generates individual gene trees, which fail to capture the complexity of species' evolutionary history (Song et al. 2012). Yet, Physcraper makes it straightforward to gather alignments and gene trees for multiple loci from a taxonomic group of interest, which can then be used by any tool that performs coalescent analyses to generate species trees, e.g., ASTRAL (Mirarab et al. 2014), BEAST2 (Bouckaert et al. 2019), SVD Quartets (Chifman & Kubatko 2014)).

- Physcraper not only links molecular and tree databases, it can also link phylogenies to various biological databases, such as GBIF, the Paleobiology Database, and others, by leveraging the integrated taxonomy from OpenTree (Rees & Cranston 2017).
- Physcraper in conjuction with OpenTree can be used to rapidly (in a matter of hours) address challenges overarching both fields of ecology and evolution, such as placing newly discovered species phylogenetically (Webb et al. 2010), obtaining trees for ecophylogenetic studies (Helmus & Ives 2012), systematizing molecular (and other) databases, i.e., curating taxonomic assignations (San Mauro & Agorreta 2010), and generating custom species trees for ecological and evolutionary downstream analyses (Stoltzfus et al. 2013).
- Data repositories hold more information than meets the eye. Besides the main data, they are rich sources of
 metadata that can be leveraged for the advantage of all areas of biology as well as the advancement of scientific
 policy and applications. Usually, initial ideas about the data are changed by new analyses. Physcraper can
 provide context for these ideas by streamlining inferences integrating new data with existing knowledge.

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- The authors have no conflict of interest to declare.

6 Authors' Contributions

- LLSR wrote the manuscript, alignment code, documentation, performed analyses and developed examples;
- ²⁴³ MK wrote code for ncbidataparser module, filtering of sequences per OTU and using offline blast searches,
- wrote documentation and tests; EJM conceived study, wrote most of the code, documentation and tests. All
- ²⁴⁵ authors contributed to the manuscript and gave final approval for publication.

²⁴⁶ 7 Data Archiving

- ²⁴⁷ Physcraper source code available at https://github.com/McTavishLab/physcraper
- ²⁴⁸ Documentation available at https://physcraper.readthedocs.io/en/latest/index.html
- Illustrated examples available at https://github.com/McTavishLab/physcraperex
- This is a reproducible manuscript available at https://github.com/McTavishLab/physcraper_ms

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