- DateLife: leveraging databases and analytical tools to reveal the dated Tree of Life
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Abstract

Date estimates for times of evolutionary divergences are key data for research in the natural 18 sciences. These estimates also provide valuable information for for education, science 19 communication and policy decisions. Although achieving a high-quality reconstruction of a 20 phylogenetic tree with branch lengths proportional to absolute time (chronogram), is a 21 difficult and time-consuming task, the increased availability of fossil and molecular data, and 22 time-efficient analytical techniques has resulted in many recent publications of large chronograms for a large number and wide diversity of organisms. When these estimates are shared in public, open databases this wealth of expertly-curated and peer-reviewed data on time of evolutionary origin is exposed in a programatic and reusable way. Intensive and localized efforts have improved data sharing practices, as well as incentivizited open science in biology. Here we present DateLife, a service implemented as an R package and an Rshiny 28 website application available at www.datelife.org/query/, that provides functionalities for efficient and easy finding, summary, reuse, and reanalysis of expert, peer-reviewed, public data on time of evolutionary origin. The main DateLife workflow constructs a chronogram 31 for any given combination of taxon names, by searching a local chronogram database 32 constructed and curated from the Open Tree of Life Phylesystem phylogenetic database, 33 which incorporates phylogenetic data from TreeBASE database as well. We implement and test methods for summarizing time data from multiple source chronograms using supertree 35 and congruification algorithms, and using age data extracted from source chronograms as secondary calibration points to add branch lengths proportional to absolute time to a tree 37 topology. DateLife will be useful to increase awereness on the existing variation in expert time of divergence data, and can foster exploration of the effect of alternative divergence time hypothesis on the results of analyses, providing a framework for a more informed interpretation of evolutionary results.

Keywords: Tree; Phylogeny; Scaling; Dating; Ages; Divergence times; Open Science;

- ⁴³ Congruification; Supertree; Calibrations; Secondary calibrations
- Word count: 4204

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46 Introduction

Chronograms –phylogenies with branch lengths proportional to time– provide key data for the study of natural processes in many areas of biological research, such as developmental biology (Delsuc et al., 2018; Laubichler & Maienschein, 2009), conservation biology (Felsenstein, 1985; C. Webb, 2000), historical biogeography (Posadas, Crisci, & Katinas, 2006), and species diversification (Magallon & Sanderson, 2001; Morlon, 2014).

Building a chronogram is not an easy task. It requires obtaining and curating data to construct a phylogeny; selecting and placing appropriate calibrations on the phylogeny using independent age data points from the fossil record or other dated events, and inferring the full dated tree. Estimating accurate chronograms generally requires specialized biological training, taxonomic domain knowledge, and a non-negligible amount of research time, computational resources and funding.

Here we present the DateLife software application, available as an R package and as an online Rshiny interactive website at www.datelife.org/query/, which captures data from published chronograms, and make these data readily accessible to users. DateLife features a versioned, open and fully public chronogram database (McTavish et al., 2015) storing age information in a computer readable format (Vos et al., 2012), an automated and programmatic way of accessing the data (Stoltzfus et al., 2013) and methods to summarize and compare age data.

Description

The DateLife algorithm is fully implemented using the R language. The latest stable version of the R package datelife is available from the CRAN repository (v0.6.2;
Sanchez-Reyes et al. (2022)), and relies on functionalities from various biological R packages:
ape (Paradis, Claude, & Strimmer, 2004), bold (Chamberlain et al., 2019), geiger (Harmon,

Weir, Brock, Glor, & Challenger, 2008), paleotree (Bapst, 2012), phyloch (Heibl, 2008),
phylocomr (Ooms & Chamberlain, 2018), phytools (Revell, 2012), rotl (Michonneau, Brown,
Winter, 2016), and taxize (Chamberlain & Szöcs, 2013; Chamberlain et al., 2019). Figure
provides a graphical summary of the three main steps of the DateLife algorithm: providing
an input, searching a chronogram database, and summarizing results from the search.

75 Providing an input

DateLife starts with an input query consisting of at least two taxon names, which can be provided as a comma separated character string, or as tip labels on a tree. If the input is a tree, it can be provided as a classic newick character string (Archie et al., 1986), or as a "phylo" R object (Paradis et al., 2004). The input tree is not required to have branch lengths, and its topology is used in the summary steps described below.

DateLife accepts scientific names as input. These names can belong to any inclusive taxonomic group (e.g., genus, family, tribe, etc.) or binomial specific. Subspecies and variants are ignored. If an input taxon name belongs to an inclusive taxonomic group the algorithm has two alternative behaviors defined by the "get species from taxon" flag. If the flag is active, the DateLife algorithm retrieves all species names within the inclusive taxonomic group and adds them to the input. If the flag is inactive, DateLife ignores the inclusive taxon names from the input.

Input scientific names are processed using a Taxonomic Name Resolution Service (TNRS), which increases the probability of correctly finding the queried taxon names in the chronogram database. TNRS detects, corrects and standardizes name misspellings and typos, variant spellings and authorities, and nomenclatural synonyms to a single taxonomic standard. DateLife implements TNRS using OpenTree's taxonomy as standard (Open Tree Of Life et al., 2016; Rees & Cranston, 2017).

The processed input taxon names are saved as an R object of a newly defined class

datelifeQuery that is used in the following steps. This object contains the processed
names, the corresponding OpenTree taxonomic id numbers, and the topology of the input
tree if any was provided.

98 Searching the database

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A DateLife search consists of matching processed taxon names to tip labels in a chronogram database. Chronograms with at least two matching tip labels are identified and pruned down to preserve only the matched tips.

Matching pruned chronograms are stored as individual patristic distance matrices
(Figure 1 subfigure X). This matrix consists of ...???? the pairwise distance between pairs
of query taxa which are in that input tree, in units of millions of years.

This format speeds up extraction of pairwise taxon ages of the queried taxa, as opposed to searching the ancestor node of a pair of taxa in a "phylo" object or newick string. The patristic matrices are also associated to the study citation where the original chronogram was published, and stored as an R object of the newly defined class datelifeResult.

DateLife's chronogram database latest version consist of 253 chronograms published in
187 different studies. It is constructed from OpenTree's phylogenetic database, the
Phylesystem, which constitutes an open source of expert phylogenetic knowledge with rich
metadata (McTavish et al., 2015) that allows automatic and reproducible construction of a
chronogram database. New chronograms can be added to Phylesystem by any user and are
immediately publicly available, and the DateLife database can be updated to include those
new data within a run.

116 Summarizing search results

At this point, summary information is extracted from the datelifeResult object to inform decisions for the subsequent steps in the user workflow. Age data from the matching

- pruned chronograms is summarized and used to generate a single summary chronogram.
- Other basic summary information available to the user is:
- 1. The matching pruned chronograms as newick strings or "phylo" objects.
- 2. The ages of the root of all matching pruned chronograms. This can correspond to the age of the most recent common ancestor (mrca) of your group of interest if the pruned chronograms have all taxa belonging to the group. If not, the root corresponds to the mrca of a subgroup withing your group of interest.
- 3. Study citations where original chronograms were published.

- 4. A report of input taxon names matches across pruned chronograms.
 - 5. The single matching pruned chronogram with the most input taxon names.
- Identifying groves.— To generate a single summary chronogram, the DateLife
 algorithm starts by identifying the matching pruned chronograms that form a grove, roughly,
 a sufficiently overlapping set of taxa between trees, by implementing definition 2.8 for
 n-overlap from Ané et al. (2009). In rare cases, a group of trees can have multiple groves. By
 default, DateLife chooses the grove with the most taxa, however, the "criterion = trees" flag
 allows the user to choose the grove with the most trees instead.
- Choosing a topology.— DateLife requires a tree topology to summarize age data upon. Users can provide one as input from the literature, or one of their own making. If no topology is provided, DateLife automatically subsets one from the OpenTree synthetic tree (Open Tree Of Life et al., 2019).
- DateLife can also reconstruct branch lengths proportional to substitution rates on a fixed tree topology using available genetic data from BOLD.
- Congruifying nodes.— DateLife then implements the congruification method (Eastman, Harmon, & Tank, 2013) to find nodes belonging to the same clade across matching pruned chronograms. Congruified node ages stored as a

congruifiedCalibrations object are then matched to nodes in the chosen tree topology and stored as a matchedCalibrations object.

Summarizing node ages.— DateLife summarizes matched calibrations into a single patristic distance matrix using different methods. Summarizing options implemented include Super Distance Matrix method (SDM, Criscuolo, Berry, Douzery, & Gascuel, 2006) and summary statistics such as median, minimum and maximum ages.

Dating the tree topology.— Summarized calibrations can be applied as secondary calibrations with different dating methods currently supported within DateLife: MrBayes (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003), PATHd8 (Britton, Anderson, Jacquet, Lundqvist, & Bremer, 2007), BLADJ (Campbell O. Webb, Ackerly, & Kembel, 2008; Campbell O. Webb & Donoghue, 2005), and treePL (Stephen A. Smith & O'Meara, 2012).

By default, DateLife implements the Branch Length Adjuster (BLADJ) algorithm that assigns ages to nodes with no data evenly between nodes with age data, which minimizes age variance in the resulting chronogram (Campbell O. Webb et al., 2008). When there is conflict in ages across node with age data, the algorithm ignores ages that are older than parent nodes and/or younger than descendant nodes.

If there is no information on the age of the root in the chronogram database, users can provide an estimate from the literature. If none is provided, DateLife assigns an arbitrary age to the root as 10% older than the oldest age available within the group.

Visualizing results.— Finally, users can save all source and summary chronograms in
formats that permit reuse and reanalyses (newick and R "phylo" format), as well as view
and compare results graphically, or construct their own graphs using datelife's chronogram
plot generation functions.

168 Benchmark

datelife's code speed was tested on an Apple iMac with one 3.4 GHz Intel Core i5 169 processor. We registered variation in computing time of query processing and search through 170 the database relative to number of queried taxon names. Query processing time increases 171 roughly linearly with number of input taxon names, and increases considerably if TNRS is 172 activated. Up to ten thousand names can be processed and searched in less than 30 minutes 173 with the most time consuming settings. Once names have been processed as described in 174 methods, a name search through the chronogram database can be performed in less than a 175 minute, even with a very large number of taxon names (Fig. ??). datelife's code 176 performance was evaluated with a set of unit tests designed and implemented with the R 177 package testthat (R Core Team, 2018) that were run both locally with the devtools package 178 (R Core Team, 2018), and on a public server -via GitHub, using the continuous integration 179 tool Travis CI (https://travis-ci.org). At present, unit tests cover more than 40% of 180 datelife's code (https://codecov.io/gh/phylotastic/datelife). 181

Case study

We illustrate the DateLife algorithm using a group within the Passeriform birds
encompassing the family of true finches, Fringillidae and allies as case study. The first
example analyses 6 bird species and shows all steps of the algorithm. The second example is
a real life application

187 Small example

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We chose 6 bird species associated to true finches at random. The sample includes two species of cardinals: the black-thighed grosbeak – *Pheucticus tibialis* and the crimson-collared grosbeak – *Rhodothraupis celaeno*; three species of buntings: the yellowhammer – *Emberiza citrinella*, the pine bunting – *Emberiza leucocephalos* and the yellow-throated bunting – *Emberiza elegans*; and one species of tanager, the vegetarian finch – *Platyspiza crassirostris*.

Processing input names found that *Emberiza elegans* is synonym for *Schoeniclus* 193 elegans in the reference taxonomy. DateLife used the processed input names to search the 194 local chronogram database and found 9 matching chronograms in 6 different studies. Three 195 studies matched five input names (Barker, Burns, Klicka, Lanyon, & Lovette, 2015; Hedges, 196 Marin, Suleski, Paymer, & Kumar, 2015; Jetz, Thomas, Joy, Hartmann, & Mooers, 2012), 197 one study matched four input names (Hooper & Price, 2017) and two studies matched two 198 input names (Barker, Burns, Klicka, Lanyon, & Lovette, 2013; Burns et al., 2014). No 199 studies matched all input names. Together, matching chronograms have 28 unique age data 200 points. All nodes have age data. As fixed tree topology, DateLife used OpenTree's synthetic 201 tree as default and mapped age data to nodes in the tree. As expected, more inclusive nodes 202 (e.g., node "n1") have more age data than less inclusive nodes (e.g., node "n5"). The 203 processing step allowed discovering five data points for node "n4" that would not have had any data otherwise. Age summary statistics per node were calculated and tested as secondary calibrations to date the tree topology using the BLADJ algorithm. Age data for node "n2" was excluded as final calibration because it is older than age data of a more inclusive node. 207

208 Real life application

A college educator wishes to obtain state-of-the-art data on time of evolutionary origin 209 of species belonging to the true finches for their class. They decide to use datelife because 210 they are teaching best practices for reproducibility. Students have the option to go to the 211 website at www.datelife.org and perform an interactive run. However, the educator also 212 wants the students to practice their R skills. The first step is to run a datelife query using the "get species from taxon" flag. This will get all recognised species names within their 214 chosen inclusive taxon. The Fringillidae has 289 species, according to the Open Tree of Life taxonomy. Once with a curated set of species taxon names, the next step is to run a 216 datelife search that will find all chronograms that contain at least two species names. The 217 algorithm proceeds to prune the trees to keep matching species names on tips only, and 218

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transform the pruned trees to pairwise distance matrices. There are 13 chronograms 219 containing at least two Fringillidae species, published in 9 different studies (Barker et al., 220 2013, 2015; Burns et al., 2014; Claramunt & Cracraft, 2015; Gibb et al., 2015; Hedges et al., 221 2015; Hooper & Price, 2017; Jetz et al., 2012; Price et al., 2014). The final step is to 222 summarize the available information using two alternative types of summary chronograms, 223 median and SDM. As explained in the "Description" section, data from source chronograms 224 is first summarised into a single distance matrix and then the available node ages are used as 225 fixed node calibrations over a consensus tree topology, to obtain a fully dated tree with the 226 program BLADJ (Fig. ??). Median summary chronograms are older and have wider 227 variation in maximum ages than chronograms obtained with SDM. ??????????????? Sav 228 some things about the results! 229

Cross-validation test

Data from source chronograms can be used to date tree topologies with no branch lengths, as well as trees with branch lengths as relative substitution rates (Figs. 5 and 6). As a form of cross validation, we took tree topologies from each input study and calibrated them using time of lineage divergence data from all other source chronograms.

In the absence of branch lengths, the ages of internal nodes were recovered with a high
precision in almost all cases (except for studies 3, and 5; Fig. 5). Maximum tree ages were
only recovered in one case (study 2; Fig. 5). We also demonstrate the usage of PATHd8
(Britton et al., 2007) as an alternative method to BLADJ. For this, we run a datelife
branch length reconstruction that searches for DNA sequence data from the Barcode of Life
Data System [BOLD; Ratnasingham and Hebert (2007)] to generate branch lengths. We
were able to successfully generate a tree with BOLD branch lengths for all of the Fringillidae
source chronograms. However, dating with PATHd8 using congruified calibrations, was only
successful in three cases (studies 3, 5, and 9, shown in Fig. 6). From these, two trees have a
different sampling than the original source chronogram, mainly because DNA BOLD data for

some species is absent from the database. ??? Node ages or maximum ages?? Maximum
ages are quite different from source chronograms, but this might be explained also by the
differences in sampling between source chronograms and BOLD trees. More examples and
code used to generate these trees were developed on an open repository that is available for
consultation and reuse at https://github.com/LunaSare/datelife examples.

250 Discussion

The main goal of datelife is to make state-of-the-art information on time of lineage divergence easily accessible for comparison, reuse, and reanalysis, to researchers in all areas of science and with all levels of expertise in the matter. It is an open service that does not require any expert biological knowledge from users –besides the names of the organisms they want to work with, for any of its functionality.

At the time of writing of this manuscript (Mar 28, 2022), datelife's database has 253 256 chronograms, pulled entirely from OpenTree's database, the Phylesystem (McTavish et al., 257 2015). A unique feature of OpenTree's Phylesystem is that the community can add new state-of-the-art chronograms any time. As chronograms are added to Phylesystem, they are incorporated into an updated datelife's database that is assigned a new version number, followed by a package release on CRAN. datelife's chronogram database is updated as new 261 chronogram data is added to Phylesystem, at a minimum of once a month and a maximum 262 of every 6 months. Users can also upload new chronograms to OpenTree themselves, and 263 trigger an update of their local datelife database to incorporate the new chronograms, to 264 have them immediately available for analysis. 265

Incorporation of more chronograms into datelife's database is crucial to improve its services. One option to increase chronogram number in the database is the Dryad data repository. Methods to automatically mine chronograms from Dryad could be designed and implemented. However, Dryad's metadata system has no information to automatically detect

branch length units, and those would still need to be determined manually by a curator.

The largest, and taxonomically broadest, summary chronogram currently available 271 from OpenTree was constructed using age data from 2,274 published chronograms (Hedges et 272 al., 2015). However the source chronograms used as input data for this tree are not available 273 in computer readable format for reuse or reanalysis. As this tree is part of datelife's 274 database, the amount of lineages that can be queried using datelife (99474 unique 275 terminal taxa) is substantial. Access to the input chronograms used to generate the Hedges 276 et al. summary tree would improve measures of uncertainty in DateLife, but they are 277 available only as image files and not as usable data (timetree.org). We would like to emphasize on the importance of sharing chronogram data for the benefit of the scientific community as a whole, into repositories that require expert input and manual curation, such 280 as OpenTree's Phylesystem (McTavish et al., 2015). 281

By default, datelife currently summarizes all source chronograms that overlap with 282 at least two species names. Users can exclude source chronograms if they have reasons to do 283 so. Strictly speaking, the best chronogram should reflect the real time of lineage divergence 284 accurately and precisely. To our knowledge, there are no good measures to determine 285 independently if a chronogram is better than another. Some measures that have been 286 proposed are the proportion of lineage sampling and the number of calibrations used 287 Magallón, Gómez-Acevedo, Sánchez-Reyes, & Hernández-Hernández (2015). Several 288 characteristics of the data used for dating analyses as well as from the output chronogram 289 itself, could be used to score quality of source chronograms. Some characteristics that are often cited in published studies as a measure of improved age estimates as compared to previously published estimates are: quality of alignment (missing data, GC content), lineage sampling (strategy and proportion), phylogenetic and dating inference method, number of 293 fossils used as calibrations, support for nodes and ages, and magnitude of confidence 294 intervals. DateLife provides an opportunity to capture concordance and conflict among date estimates, which can also be used as a metric for chronogram reliability.

Scientists usually also favor chronograms constructed using primary calibrations (ages 297 obtained from the fossil or geological record) to ones constructed with secondary calibrations 298 (ages coming from other chronograms) (Schenk, 2016). It has been observed with simulations 299 that divergence times inferred with secondary calibrations are significantly younger than 300 those inferred with primary calibrations in analyses performed with Bayesian inference 301 methods when priors are implemented in similar ways in both analyses (Schenk, 2016). 302 However, secondary calibrations can be applied using other dating methods that do not 303 require setting priors, such as penalized likelihood (Sanderson, 2003), or as fixed ages, 304 potentially mitigating the bias reported with Bayesian methods. Certainly, further studies 305 are required to fully understand the effect of using secondary calibrations on time estimates 306 and downstream analyses. 307

Furthermore, even chronograms obtained with primary fossil data can vary 308 substantially in time estimates between lineages, as observed from the comparison of source 300 chronograms in the Fringillidae example. This observation is often encountered in the 310 literature (see, for example, the ongoing debate about crown group age of angiosperms 311 (Barba-Montoya, Reis, Schneider, Donoghue, & Yang, 2018; Magallón et al., 2015; Ramshaw 312 et al., 1972; Sanderson & Doyle, 2001; Sauquet, Ramírez-Barahona, & Magallón, 2021). For 313 some studies, especially ones based on branch lengths (e.g., studies of species diversification, timing of evolutionary events, phenotypic trait evolution), using a different chronogram may return different results (Title & Rabosky, 2016). Stitching together these chronograms can 316 create a larger tree that uses information from multiple studies, but the effect of 317 uncertainties and errors at this level on downstream analyses is still largely unknown. 318

Summarizing chronograms might also imply summarizing fundamentally distinct evolutionary hypotheses. For example, two different researchers working on the same clade both carefully select and argument their choices of fossil calibrations. Still, if one researcher

decides a fossil will calibrate the ingroup of a clade, while another researcher uses the same one to calibrate outside the clade, the resulting age estimates will often differ substantially, 323 as the placement of calibrations as stem or crown group is proved to deeply affect estimated 324 times of lineage divergence (Sauquet, 2013). Trying to summarize the resulting chronograms 325 into a single one using simple summary statistics can erase many types of relevant 326 information from the source chronograms. Accordingly, the prevailing view is that we should 327 favor time of lineage divergence estimates obtained from a single analysis, using fossil data as 328 primary sources of calibrations, and using fossils that have been widely discussed and 329 curated as calibrations to date other trees, making sure that all data used in the analysis 330 reflect a coherent evolutionary history (Antonelli et al., 2017). However, the exercise of 331 summarizing different chronograms has the potential to help getting a single global 332 evolutionary history for a lineage by putting together evidence from different hypothesis. Choosing the elements of the chronograms that we are going to keep and the ones that we 334 are going to discard is key, since we are potentially loosing important parts of the 335 evolutionary history of a lineage that might only be reflected in source chronograms and not 336 on the summary chronogram (Sauguet et al., 2021). 337

Nonetheless, in ecology and conservation biology, incorporating at least some data on 338 lineage divergence times represents a relevant improvement for testing alternative hypothesis 339 using phylogenetic distance (Campbell O. Webb et al., 2008). Hence, we integrated into 340 datelife's workflow different ways of estimating node ages in the absence of calibrations and 341 branch length information for taxa lacking this information. "Making up" branch lengths is 342 an accepted practice in scientific publications: Jetz et al. (2012), created a time-calibrated tree of all 9,993 bird species, where 67% had molecular data and the rest was simulated; Rabosky et al. (2018) created a time-calibrated tree of 31,536 ray-finned fishes, of which only 37% had molecular data; Stephen A. Smith and Brown (2018) constructed a tree of 353,185 seed plants where only 23% had molecular data. Obviously, there are risks in this practice! 347 Taken to the extreme, one could make a fully resolved, calibrated tree of all modern and

extinct taxa using a single taxonomy and a single calibration with the polytomy resolution
and branch estimation methods. There has yet to be a thorough analysis of what can go
wrong when one extends inferences beyond the data in this way, so we urge caution; we also
urge readers to follow the example of many of the large tree papers cited above and make
carefully consider the statistical assumptions being made, and assess the consistency of the
results with prior work.

355 Conclusions

Divergence time information is key to many areas of evolutionary studies: trait
evolution, diversification, biogeography, macroecology and more. It is also crucial for science
communication and education, but generating chronograms is difficult, especially for those
who want to use phylogenies but who are not systematists, or do not have the time to
acquire and develop the necessary knowledge and data curation skills. Moreover, years of
primarily public funded research have resulted in vast amounts of chronograms that are
already available on scientific publications, but hidden to the public and scientific community
for reuse.

The datelife R package allows easy and fast summarization of publicly available information on time of lineage divergence. This provides a straightforward way to get an informed idea on the state of knowledge of the time frame of evolution of different regions of the tree of life, and allows identification of regions that require more research or that have conflicting information. It is available as an R package, or a web-based R shiny app at dates.opentreeloflife.org/datelife. Both summary and newly generated trees are useful to evaluate evolutionary hypotheses in different areas of research. The DateLife project helps with awareness of the existing variation in expert time of divergence data, and will foster exploration of the effect of alternative divergence time hypothesis on the results of analyses, nurturing a culture of more cautious interpretation of evolutionary results.

Availability

datelife is free and open source and it can be used through its current website 375 http://www.datelife.org/query/, through its R package, and through Phylotastic's project 376 web portal http://phylo.cs.nmsu.edu:3000/. datelife's website is maintained using 377 RStudio's shiny server and the shiny package open infrastructure, as well as Docker. 378 datelife's R package stable version is available for installation from the CRAN repository 379 (https://cran.r-project.org/package=datelife) using the command install.packages(pkgs 380 = "datelife") from within R. Development versions are available from the GitHub 381 repository (https://github.com/phylotastic/datelife) and can be installed using the 382 command devtools::install github("phylotastic/datelife"). 383

Supplementary Material

Code used to generate all versions of this manuscript, the biological examples, as well as the benchmark of functionalities are available at datelifeMS1, datelife_examples, and datelife_benchmark repositories in LLSR's GitHub account.

Funding Funding

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393

Funding was provided by the US National Science Foundation (NSF) grants
ABI-1458603 to Datelife project and DBI-0905606 to the National Evolutionary Synthesis
Center (NESCent), and the Phylotastic project Grant ABI-1458572, and the OpenTree grant
ABI-1759846.

Acknowledgements

The DateLife project was born as a prototype tool aiming to provide these services, and was developed over a series of hackathons at the National Evolutionary Synthesis Center, NC, USA (Stoltzfus et al., 2013). We thank colleagues from the O'Meara Lab at the University of Tennesse Knoxville for suggestions, discussions and software testing. The late

National Evolutionary Synthesis Center (NESCent), which sponsored hackathons that led to 398 initial work on this project. The team that assembled datelife's first proof of concept: 399 Tracy Heath, Jonathan Eastman, Peter Midford, Joseph Brown, Matt Pennell, Mike Alfaro, 400 and Luke Harmon. The Open Tree of Life project that provides the open, metadata rich 401 repository of trees used for datelife. The many scientists who publish their chronograms in 402 an open, reusable form, and the scientists who curate them for deposition in the Open Tree 403 of Life repository. The NSF for funding nearly all the above, in addition to the ABI grant 404 that funded this project itself. 405

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to be formatted in the same way as the general text (double spaced and linenumbered)

LTT plots showing results from the cross-validation analyses of trees with branch length reconstructed with data from the Barcode of Life Database (BOLD) dated using PATHd8. We could construct a tree with branch lengths for all source chronograms.

However, dating with PATHd8 was only successful in three source chronograms shown here.

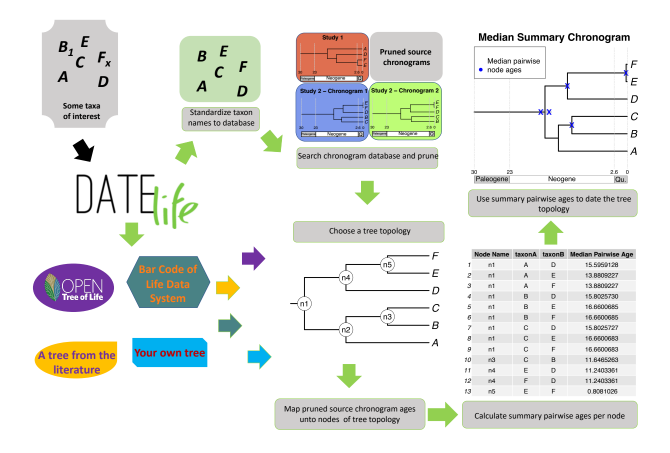


FIGURE 1. Stylized DateLife workflow. This shows the general worflows and analyses that can be performed with datelife, via the R package or through the website at http://www.datelife.org/. Details on the functions involved on each workflow are shown in datelife's R package vignette.

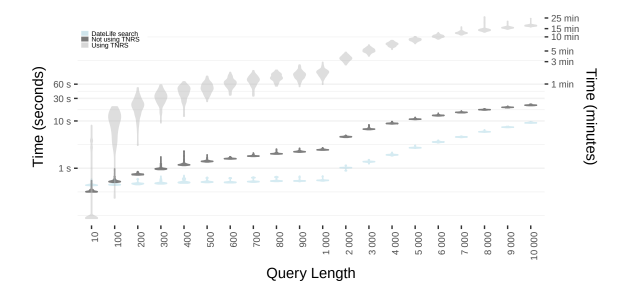


FIGURE 2. Computation time of query processing and search across datelife's chronogram database relative to number of input taxon names. We sampled N names from the class Aves for each cohort 100 times and then performed a search with query processing not using the Taxon Names Resoultion Service (TNRS; dark gray), and using TNRS (light gray). We also performed a search using the already processed query for comparison (light blue).

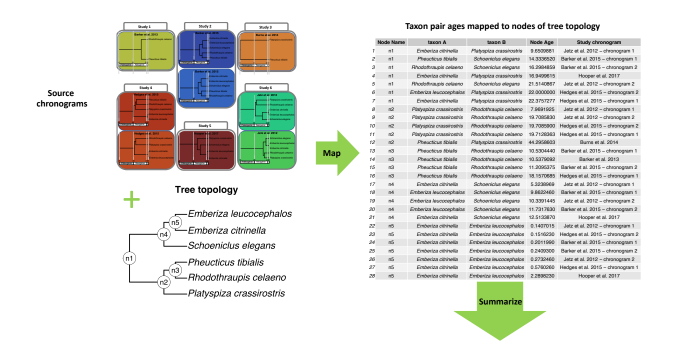


FIGURE 3. Age data results of a DateLife search of a small sample of 6 bird species within the Passeriformes. Input names were found across 9 chronograms within 6 independent studies (Barker et al. (2012), Barker et al. (2015), Burns et al. (2014), Hedges et al. (2015), Hooper and Price (2017), Jetz et al. (2012).) This revealed 28 age data points for the queried species names.

Summary	of of	map	ped	taxon	pair	age	data
	•				P		

	Node Name	taxon A	taxon B	Pairwise Median Age	Node Median Age
1		Pheucticus tibialis	Emberiza citrinella	16.298486	
2		Pheucticus tibialis	Emberiza leucocephalos	16.298486	
3		Platyspiza crassirostris	Emberiza citrinella	21.514085	
4		Platyspiza crassirostris	Emberiza leucocephalos	21.514085	
5	n1	Rhodothraupis celaeno	Emberiza citrinella	20.408031	19.301977
6		Rhodothraupis celaeno	Emberiza leucocephalos	20.408031	
7		Schoeniclus elegans	Pheucticus tibialis	15.316069	
8		Schoeniclus elegans	Platyspiza crassirostris	19.301977	
9		Schoeniclus elegans	Rhodothraupis celaeno	17.800231	
10	n2	Platyspiza crassirostris	Pheucticus tibialis	32.004348	25.856467327225
11		Rhodothraupis celaeno	Platyspiza crassirostris	19.708587	
12	n3	Rhodothraupis celaeno	Pheucticus tibialis	10.873723	10.87372335475
13	n4	Schoeniclus elegans	Emberiza citrinella	10.647794	10.6477935
14		Schoeniclus elegans	Emberiza leucocephalos	10.647794	
15	n5	Emberiza leucocephalos	Emberiza citrinella	0.273246	0.273246

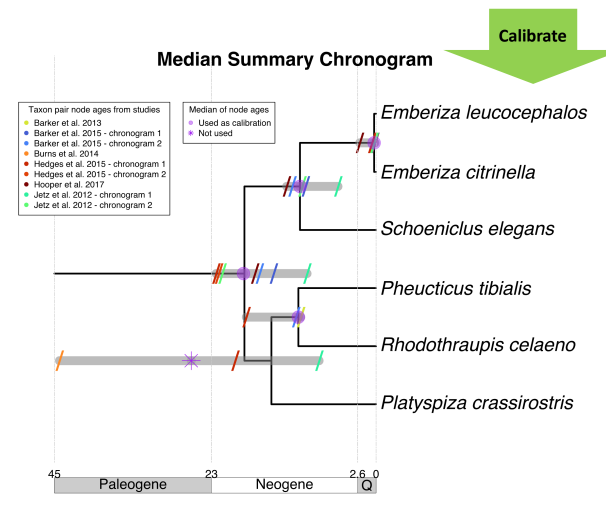


FIGURE 4. Summarized age data is used as secondary calibrations to date a tree topology as a summary chronogram.

- Fringillidae source chronograms used as calibrations
 Fringillidae source chronograms used only as topology
 Fringillidae new chronograms generated with PATHd8

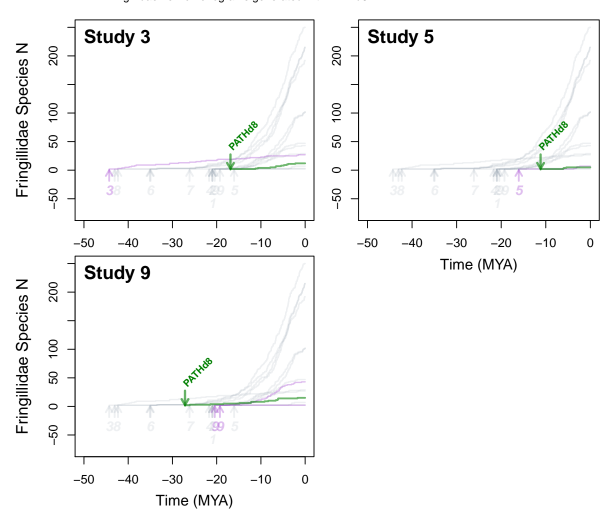


Figure 6