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

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Chronograms –phylogenies with branch lengths proportional to time– represent key 19 data on timing of evolutionary events for the study of natural processes in many areas of biological research. Chronograms also provide valuable information that can be used for education, science communication, and conservation policy decisions. Yet, achieving a high-quality reconstruction of a chronogram is a difficult and resource-consuming task. Here we present DateLife, a phylogenetic software implemented as an R package and an R Shiny web application available at www.datelife.org, that provides services for efficient and easy 25 discovery, summary, reuse, and reanalysis of node age data mined from a curated database of 26 expert, peer-reviewed, and openly available chronograms. The main DateLife workflow starts 27 with one or more scientific taxon names provided by a user. Names are processed and 28 standardized to a unified taxonomy, allowing DateLife to run a name match across its local chronogram database that is curated from Open Tree of Life's phylogenetic repository, and extract all chronograms that contain at least two queried taxon names, along with their 31 metadata. Finally, node ages from matching chronograms are mapped using the congruification algorithm to corresponding nodes on a tree topology, either extracted from Open Tree of Life's synthetic phylogeny or one provided by the user. Congruified node ages are used as secondary calibrations to date the chosen topology, with or without initial 35 branch lengths, using different phylogenetic dating methods such as BLADJ, treePL, PATHd8 and MrBayes. We performed a cross-validation test to compare node ages resulting from a DateLife analysis (i.e., phylogenetic dating using secondary calibrations) to those from the original chronograms (i.e., obtained with primary calibrations), and found that DateLife's node age estimates are consistent with the age estimates from the original chronograms, with the largest variation in ages occurring around topologically deeper nodes. Because the results from any software for scientific analysis can only be as good as the data used as input,

- 43 we highlight the importance of considering the results of a DateLife analysis in the context of
- the input chronograms. DateLife can help to increase awareness of the existing disparities
- 45 among alternative hypotheses of dates for the same diversification events, and to support
- exploration of the effect of alternative chronogram hypotheses on downstream analyses,
- 47 providing a framework for a more informed interpretation of evolutionary results.
- 48 Keywords: Tree; Phylogeny; Scaling; Dating; Ages; Divergence times; Open Science;
- Congruification; Supertree; Calibrations; Secondary calibrations.
- 50 Word count: 6817

Chronograms –phylogenies with branch lengths proportional to time– provide key data on evolutionary time frame for the study of natural processes in many areas of biological research, such as comparative analysis (Freckleton, Harvey, & Pagel, 2002; Harvey, Pagel, & others, 1991), developmental biology (Delsuc et al., 2018; Laubichler & Maienschein, 2009), conservation biology and ecology (Felsenstein, 1985; 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 a
homology hypothesis 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 a full dated tree. All of this entails specialized biological training,
taxonomic domain knowledge, and a significant amount of research time, computational
resources and funding.

Here we present the DateLife project which has the main goal of extracting and
exposing age data from published chronograms, making age data readily accessible to a
wider community for reuse and reanalysis in research, teaching, science communication and
conservation policy. DateLife's core software application is available as an R package
(Sanchez-Reyes et al., 2022), and as an online Rshiny interactive website at www.datelife.org.
It features key elements for scientific reproducibility, such as a curated, versioned, open and
fully public chronogram database (McTavish et al., 2015) that stores data in a
computer-readable format (Vos et al., 2012); automated and programmatic ways of accessing
and downloading the data, also in a computer-readable format (Stoltzfus et al., 2013); and
methods to summarize and compare the data.

Description

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DateLife's core software applications are implemented in the R package datelife, and

relies on functionalities from other biological R packages: ape (Paradis, Claude, & Strimmer, 2004), bold (Chamberlain, 2018), geiger (Pennell et al., 2014), msa (Bodenhofer, Bonatesta, Horejš-Kainrath, & Hochreiter, 2015), paleotree (Bapst, 2012), phyloch (Heibl, 2008), phylocomr (Ooms & Chamberlain, 2018), phytools (Revell, 2012), rotl (Michonneau, Brown, & Winter, 2016), and taxize (Chamberlain, 2018; Chamberlain & Szöcs, 2013). There are three main steps to the DateLife workflow: 1) creating a search query, 2) searching a database, and 3) summarizing results from the search.

Creating a Search Query

DateLife starts by processing an input consisting of the scientific name of at least one taxon. Multiple input names 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 in the next section.

DateLife processes input scientific names using a Taxonomic Name Resolution Service 90 (TNRS), which increases the probability of correctly finding the queried taxon names in the 91 chronogram database. TNRS detects, corrects and standardizes name misspellings and typos, variant spellings and authorities, and nomenclatural synonyms to a single taxonomic standard (Boyle et al., 2013). TNRS also allows to correctly choose between homonyms, by considering other taxa provided as input to infer the taxonomic context of the homonym. DateLife implements TNRS using the Open Tree of Life (OpenTree) unified Taxonomy (OTT, Open Tree Of Life et al., 2016; Rees & Cranston, 2017) as standard, storing taxonomic identification numbers (OTT ids) for further processing and analysis. Other taxonomies currently supported by DateLife are the National Center of Biotechnology Information (NCBI) taxonomic database (Schoch et al., 2020), the Global Biodiversity 100 Information Facility (GBIF) taxonomic backbone (GBIF Secretariat, 2022), and the Interim 101

Register of Marine and Non-marine Genera (IRMNG) database (Rees et al., 2017).

Besides binomial species names, DateLife accepts scientific names from any inclusive 103 taxonomic group (e.g., genus, family, tribe), as well as subspecific taxonomic variants (e.g., 104 subspecies, variants, strains). If a taxon name belongs to an inclusive taxonomic group, 105 DateLife has two alternative behaviors defined by the "get species from taxon" flag. If the 106 flag is active, DateLife retrieves all species names within a taxonomic group provided, from a 107 standard taxonomy of choice, and adds them to the search query. In this case, subspecific 108 variants are excluded. If the flag is inactive, DateLife excludes inclusive taxon names from 109 the search query, and species and subspecific variant names are processed as provided by the 110 user. The processed taxon names are saved as an R object of a newly defined class, 111 datelifeQuery, that is used in the following steps. This object contains the input names 112 standardized to a taxonomy of choice (OTT by default), the corresponding OTT id numbers, 113 and the topology of an input tree, if one was provided. 114

Searching a Chronogram Database

At the time of writing of this manuscript (Nov 07, 2023), DateLife's chronogram
database latest version consist of 253 chronograms published in 187 different studies. It is
curated from OpenTree's phylogenetic database, the Phylesystem, which constitutes an open
source of expert and peer-reviewed phylogenetic knowledge with rich metadata (McTavish et
al., 2015), which allows automatic and reproducible assembly of our chronogram database.
Datelife's chronogram database is navigable as an R data object within the datelife R
package.

A unique feature of the Phylesystem is that any user can add new published,

state-of-the-art chronograms any time, through OpenTree's curator application

(https://tree.opentreeoflife.org/curator). As chronograms are added to Phylesystem, they

can be incorporated into the chronogram database of the datelife R package, which is

currently manually updated as new chronogram data is added to Phylesystem. The updated
database is assigned a new version number, followed by a package release on CRAN. Users
can directly run datelife functions to trigger an update of their local chronogram database,
to incorporate any new chronograms to their DateLife analysis before an official database
update is released on CRAN.

A DateLife search is implemented by matching processed taxon names provided by the 132 user to tip labels in the chronogram database. Chronograms with at least two matching 133 taxon names on their tip labels are identified and pruned down to preserve only the matched 134 taxa. These matching pruned chronograms are referred to as source chronograms. Total 135 distance in units of million years (Myr) between taxon pairs within each source chronogram 136 are stored as a patristic distance matrix. The matrix format speeds up extraction of pairwise 137 taxon ages of any queried taxa, as opposed to searching the ancestor node of a pair of taxa 138 in a "phylo" object or newick string. Finally, the patristic matrices are associated to the 139 study citation where the original chronogram was published, and stored as an R object of the newly defined class datelifeResult. 141

Summarizing Search Results

Summary information is extracted from the datelifeResult object to inform decisions for subsequent steps in the analysis workflow. Basic summary information available to the user includes:

- 1. The matching pruned chronograms as newick strings or "phylo" objects.
- 2. The ages of the root of all source chronograms. These ages can correspond to the age
 of the most recent common ancestor (mrca) of the user's group of interest if the source
 chronograms have all taxa belonging to the group. If not, the root corresponds to the
 mrca of a subgroup within the group of interest.
 - 3. Study citations where original chronograms were published.

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- 4. A report of input taxon names matches across source chronograms.
 - 5. The source chronogram(s) with the most input taxon names.
- 6. Various single summary chronograms resulting from summarizing age data, generated using the methodology described next.

Choosing a Topology

DateLife requires a tree topology to summarize age data upon. We recommend that 157 users provide as input a tree topology from the literature, or one of their own making. If no 158 topology is provided, DateLife automatically extracts one from the OpenTree synthetic tree, 159 a phylogeny currently encompassing 2.3 million taxa across all life, assembled from 1,239 160 published phylogenetic trees and OpenTree's unified Taxonomy, OTT (Open Tree Of Life et 161 al., 2019). Alternatively, DateLife can combine topologies from source chronograms using a 162 supertree approach (Criscuolo, Berry, Douzery, & Gascuel, 2006). To do this, DateLife first 163 identifies the source chronograms that form a grove, roughly, a sufficiently overlapping set of 164 taxa between trees, by implementing definition 2.8 for n-overlap from Ané et al. (2009). If 165 the source chronograms do not form a grove, the supertree reconstruction will fail. In rare 166 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. The result is a single summary (i.e., supertree) topology, that combines topologies from source chronograms in a grove.

Applying Secondary Calibrations

Once a topology is chosen, DateLife applies the congruification method (Eastman,
Harmon, & Tank, 2013) that find nodes belonging to the same clade across source
chronograms, and then extracts the corresponding node ages from patristic distance matrices
stored as a datelifeResult object. Note that by definition, these matrices store total
distance (time from tip to tip), assuming that the terminal taxa are coeval and occur at the

present. Hence, node ages correspond to half the values stored in the datelifeResult matrices. A table of congruified node ages that can be used as calibrations for a dating analysis is stored as a congruifiedCalibrations object.

For each congruent node, the pairwise distances that traverse that node are summarized 180 into a single summary matrix using classic summary statistics (i.e., mean, median, minimum 181 and maximum ages), and the Supermatrix Distance Method (SDM: Criscuolo et al., 2006), 182 which deforms patristic distance matrices by minimizing variance and then averaging them. 183 These single summary taxon pair age matrices are stored as summarized calibrations that 184 can be used as secondary calibrations to date a tree topology - with or without initial branch 185 lengths, using phylogenetic dating methods currently supported within DateLife: BLADJ 186 (Webb, Ackerly, & Kembel, 2008; Webb & Donoghue, 2005), MrBayes (Huelsenbeck & 187 Ronquist, 2001; Ronquist & Huelsenbeck, 2003), PATHd8 (Britton, Anderson, Jacquet, 188 Lundqvist, & Bremer, 2007), and treePL (Smith & O'Meara, 2012).

Dating a Tree Topology

Dating a tree without branch lengths. To date a tree topology when initial 191 branch lengths are unavailable, DateLife implements the Branch Length Adjuster (BLADJ) 192 algorithm (Webb et al., 2008; Webb & Donoghue, 2005), which only requires a tree topology 193 with no branch lengths and at least two node ages to use as calibrations, one for the tree root 194 and one for any internal node of the topology. The BLADJ algorithm fixes ages for nodes 195 with calibration data upon the given tree topology. Then, it assigns ages to nodes with no 196 available age information by distributing time evenly between calibrated nodes, minimizing age variance in the resulting chronogram. This approach has proven useful for ecological 198 analyses that require a phylogenetic time context (Webb et al., 2008). When there is conflict 199 between ages of calibrated nodes, BLADJ ignores node ages that are older than the age of a 200 parent node. The BLADJ algorithm requires a root age to run. Users can provide an 201 appropriate root age estimate of their own or one obtained from the literature. If a root age 202

is not provided and there is no information on the age of the root in the chronogram
database, DateLife chooses a random age for the root, so that a dated tree topology can be
generated with BLADJ. In this case, DateLife will provide a conspicuous warning message,
so that users are aware that the root of the chronogram was chosen at random because there
was no information available for it in the chronogram database, along with suggestions on
how the user can find and provide an appropriate age for the root of the initial topology.

An alternative to BLADJ to date tree topologies in the absence of initial branch 209 lengths that is common practice in the literature is to use a birth-death model to draw 210 branch lengths (Jetz, Thomas, Joy, Hartmann, & Mooers, 2012; Rabosky et al., 2018; Smith & Brown, 2018). In addition to the initial tree topology and nodes with age data, these methods require initial values of speciation and extinction rate parameters provided by the 213 user. DateLife implements this approach with MrBayes (Huelsenbeck & Ronquist, 2001; 214 Ronquist & Huelsenbeck, 2003), using nodes with published age data as calibration priors on 215 nodes of a tree topology with no branch lengths, a simple birth-death model with speciation 216 and extinction rate parameters that are provided by the user, and no genetic data. However, 217 BLADJ is the default option in DateLife, as it does not require any information on 218 diversification rates for the phylogenetic sample to draw from a branch length distribution. 219

Dating a tree with branch lengths.— Relative branch lengths can provide key information for phylogenetic dating, specifically for nodes without any calibration data available. While using initial branch length data is the golden standard for phylogenetic dating analyses, estimating trees with branch lengths proportional to substitution rates per site requires obtaining primary data, assembling and curating a homology (orthology) hypothesis, and choosing and implementing a method for phylogenetic inference. DateLife implements a workflow to streamline this process by applying open data from the Barcode of Life Data System, BOLD (Ratnasingham & Hebert, 2007) to obtain genetic markers for input taxa. By default, BOLD genetic sequences are aligned with MUSCLE (Edgar, 2004)

using functions from the msa R package (Bodenhofer et al., 2015). Alternatively, sequences 229 can be aligned with MAFFT (Katoh, Asimenos, & Toh, 2009), using functions from the ape 230 R package (Paradis et al., 2004). The BOLD sequence alignment is then used to obtain 231 initial branch lengths with the accelerated transformation (ACCTRAN) parsimony 232 algorithm, which resolves ambiguous character optimization by assigning changes along 233 branches of the tree as close to the root as possible (Agnarsson & Miller, 2008), resulting in 234 older internal nodes as compared to other parsimony algorithms (Forest et al., 2005). The 235 parsimony branch lengths are then optimized using Maximum Likelihood, given the 236 alignment, the topology and a simple Jukes-Cantor model, producing a BOLD tree with 237 branch lengths proportional to expected number of substitutions per site. Both parsimony 238 and ML optimizations are done with functions from the phangorn package (Schliep, 2011). 239 Due to the computing load it requires, the BOLD workflow is currently only supported through DateLife's R package. It is not yet available through the web application.

Phylogenetic dating methods supported in DateLife that incorporate branch length information from the input topology in combination with the secondary calibrations include:
PATHd8, a non-clock, rate-smoothing method to date trees (Britton et al., 2007); treePL (Smith & O'Meara, 2012), a semi-parametric, rate-smoothing, penalized likelihood dating method (Sanderson, 2002); and MrBayes (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003), a Bayesian inference program implementing Markov chain Monte Carlo (MCMC) methods to estimate a posterior distribution of model parameters.

Visualizing Results

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Finally, users can save all source and summary chronograms in formats allowing for reuse and reanalysis, such as newick and the R "phylo" format. Input and summary chronograms can be visualized and compared graphically, and users can construct their own graphs using DateLife's chronogram plot generation functions available from the R package datelifeplot (Sanchez-Reyes & O'Meara, 2022).

BENCHMARK 255

R package datelife code speed was tested on an Apple iMac with one 3.4 GHz Intel 256 Core is processor. We registered variation in computing time of query processing and search 257 through the database relative to number of queried taxon names. Query processing time 258 increases roughly linearly with number of input taxon names, and increases considerably if 250 Taxonomic Name Resolution Service (TNRS) is activated. Up to ten thousand names can be 260 processed and searched in less than 30 minutes with the most time consuming settings. Once 261 names have been processed as described in methods, a name search through the chronogram 262 database can be performed in less than a minute, even with a very large number of taxon 263 names (Fig. 1). 264

datelife's code performance was evaluated with a set of unit tests designed and 265 implemented with the R package testthat (R Core Team, 2018) that were run both locally with the devtools package (R Core Team, 2018), and on a public server using the continuous integration tool of GitHub actions (https://docs.github.com/en/actions). At present, unit 268 tests cover more than 40% of datelife's code (https://codecov.io/gh/phylotastic/datelife). Unit testing helps identify potential issues as code is updated or, more critically, as services code relies upon may change.

Case Studies

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We illustrate the DateLife workflow using a family within the passeriform birds 273 encompassing the true finches, Fringillidae, as case study. On a small example, we analysed 6 bird species, and results from each step of the workflow are shown in Figure 2. As a second 275 example, we analysed 289 bird species in the family Fringillidae that are included in the NCBI taxonomy. One clade from the full summary chronogram result from the DateLife analysis is shown Figure 3. The full chronogram for all 289 species and the results from 278 previous steps of the workflow are available as Supplementary Figures.

A Small Example

Creating a search query.— We chose 6 bird species within the Passeriformes. The 281 sample includes two species of cardinals: the black-thighed grosbeak – Pheucticus tibialis 282 and the crimson-collared grosbeak – Rhodothraupis celaeno; three species of buntings: the 283 yellowhammer – Emberiza citrinella, the pine bunting – Emberiza leucocephalos and the 284 yellow-throated bunting – Emberiza elegans; and one species of tanager, the vegetarian finch – 285 Platyspiza crassirostris. Processing of input names found that Emberiza elegans is synonym 286 for Schoeniclus elegans in the default reference taxonomy (OTT v3.3, June 1, 2021). For a 287 detailed discussion on the state of the synonym, refer to Avibase (Avibase, 2022; Lepage, 288 2004; Lepage, Vaidya, & Guralnick, 2014). Discovering this synonym allowed assigning five 280 age data points for the parent node of *Emberiza elegans*, shown as *Schoeniclus elegans* in 290 Figure 2, which would not have had any data otherwise. 291

Searching the database.— DateLife used the processed input names to search the local chronogram database and found 9 matching chronograms from 6 different studies (Fig. 2c). Three studies matched five input names (Barker, Burns, Klicka, Lanyon, & Lovette, 2015; Hedges, Marin, Suleski, Paymer, & Kumar, 2015; Jetz et al., 2012), one study matched four input names (Hooper & Price, 2017) and two studies matched two input names (Barker, Burns, Klicka, Lanyon, & Lovette, 2013; Burns et al., 2014). No studies matched all input names. Together, source chronograms provide 28 unique age data points, covering all nodes on our chosen tree topology to date (Table 1).

Summarizing search results.— DateLife obtained OpenTree's synthetic tree topology for these taxa (Fig. 2d), and congruified and mapped age data to nodes in this chosen topology, shown in Table 1. The name processing step allowed including five data points for node "n4" (parent of Schoeniclus elegans) that would not have had any data otherwise due to name mismatch. Age summary statistics per node were calculated (Table 2) and used as calibrations to date the tree topology using the BLADJ algorithm. As expected,

more inclusive nodes (e.g., node "n1") have more variance in age data than less inclusive nodes (e.g., node "n5"). Median summary age data for node "n2" was excluded as final calibration because it is older than the median age of a more inclusive node, "n1" (Fig. 2g).

An Example with the Family of True Finches

Creating a query. To obtain ages for all species within the family of true finches, 310 Fringillidae, we ran a DateLife query using the "get species from taxon" flag, which gets all 311 recognized species names within a named group from a taxonomy of choice. Following the 312 NCBI taxonomy, our DateLife query has 289 Fringillidae species names. This 313 taxon-constrained approach implies that the full DateLife analysis will be performed using a 314 tree topology and ages available for species names from a given taxonomic group, which do 315 not necessarily correspond to a monophyletic group. Users can change this behavior by 316 providing all species names corresponding to a monophyletic group as input for a DateLife 317 search, or a monophyletic tree to construct a DateLife summary. 318

Searching the database. Next, we used the processed species names in our 319 DateLife query to identify chronograms with at least two Fringillidae species as tip taxa. 320 The DateLife search identified 19 chronograms matching this criteria, published in 13 321 different studies (Barker et al., 2013, 2015; Burns et al., 2014; Claramunt & Cracraft, 2015; 322 Gibb et al., 2015; Hedges et al., 2015; Hooper & Price, 2017; Jetz et al., 2012; Kimball et al., 323 2019; Oliveros et al., 2019; Price et al., 2014; Roquet, Lavergne, & Thuiller, 2014; Uyeda, 324 Pennell, Miller, Maia, & McClain, 2017). Once identified, DateLife pruned these matching 325 chronograms to remove tips that do not belong to the queried taxon names, and transformed these pruned chronograms to pairwise distance matrices, revealing 1,206 different age data points available for species within the Fringillidae (Supplementary Table S1).

Summarizing search results.— The final step entailed congruifying and
summarizing the age data available for the Fringillidae species into two single summary
chronograms, using two different types of summary ages, median and SDM. As explained in
the "Description" section, a tree topology to summarize age data upon is required. By
default, DateLife uses the topology from OpenTree's synthetic tree that contains all taxa
from the search query. According to OpenTree's synthetic tree, species belonging to the

family Fringillidae do not form a monophyletic group (Supplementary Fig. S1). Hence, a topology containing only the 289 species from the original query was extracted from Open Tree of Life's synthetic tree v12.3 (Supplementary Fig. S2; Open Tree Of Life et al., 2019).

All 19 source chronograms (Supplementary Figs. S5-S23) were congruified to 338 OpenTree's topology shown in Supplementary Figure S2, reducing the original 1,206 node 339 age data set to 818 different data points (Supplementary Table S2) that could be used as 340 calibrations for the chosen topology (Supplementary Fig. S2). The congruent node age data 341 points were summarized for each node, resulting in 194 summary node ages. From these 21 342 were excluded as secondary calibrations because they were older than the ancestral node. 343 The remaining 173 summary node ages were used as secondary calibrations to obtain a fully 344 dated (and resolved) phylogeny with the program BLADJ (Supplementary Figure S3). 345 Results for a subgroup are shown in Figure 3.

CROSS-VALIDATION TEST

We performed a cross validation test of a DateLife analysis using the Fringillidae 348 source chronograms obtained above (Supplementary Figs. S5-S23). As inputs for a DateLife 349 analysis, we used all individual tree topologies from each of the 19 source chronograms from 350 13 studies, treating their node ages as unknown. We congruified node ages extracted from 351 chronograms from all other studies upon the individual topologies, effectively excluding 352 original ages from each topology. Finally, average node ages per node were applied as 353 secondary calibrations and smoothed with the BLADJ algorithm. We found that node ages 354 from the original studies, and ages estimated using all other age data available are generally correlated (Fig. 4). For five studies, DateLife tended to underestimate ages for topologically deeper nodes (those with many descendant taxa, aka "closer to the root") relative to the 357 original estimate, and overestimate ages for nodes closer to the tips. Accordingly, root ages 358 are generally older in the original study than estimated using cross-validated ages 359 (Supplementary Fig. S4). In general, topologically deeper nodes display the largest age

variation between node ages from the original chronograms and ages summarized with

DateLife.

363 DISCUSSION

DateLife's goal is to improve availability and accessibility of state-of-the-art data on
evolutionary time frame of organisms, to allow users from all areas of science and with all
levels of expertise to compare, reuse, and reanalyse expert age data for their own
applications. As such, it is designed as an open service that does not require any expert
biological knowledge –besides the scientific names of the species or group users want to work
with, to use any of its functionalities.

A total of 99,474 unique terminal taxa are represented in DateLife's database. 370 Incorporation of more chronograms into the database will continue to improve DateLife's 371 services. One option to increase the number of chronograms in the DateLife database is the 372 Dryad data repository. Methods to automatically mine chronograms from Dryad could be 373 designed and implemented. However, Dryad's metadata system has no information to 374 automatically detect branch length units, and those would still need to be determined 375 manually by a human curator. We would like to emphasize on the importance of sharing 376 chronogram data, including systematically curated metadata, into open repositories, such as 377 OpenTree's Phylesystem (McTavish et al., 2015) for the benefit of the scientific community as a whole. 379

As we envision that DateLife will have many interesting applications in research and
beyond, we emphasize that DateLife's results –as well as any insights gleaned from them,
largely depend on the quality of the source chronograms: low quality chronograms will
produce low quality results. The "garbage in, garbage out" problem has long been recognised
in supertree methods for summarizing phylogenetic trees (Bininda-Emonds et al., 2004). We
note that this is a surfacing issue of any automated tool for biological data analysis. For

example, DNA riddled with sequencing errors will produce generally poor alignments that
will return biased evolutionary hypothesis, independently of the quality of the analysis
software used. Again, we urge readers and DateLife users to explore all input chronograms
before using a summary chronogram resulting from a DateLife workflow.

Finally, uncertainty and variability of chronogram node age estimates might pose larger issues in some research areas than others. For example, in ecological and conservation biology studies, it has been shown that incorporating some chronogram data provides better results than when not using any age data at all, even if the node ages are not good quality (Webb et al., 2008). In the following sections we discuss the particularities of divergence times from DateLife's summary chronograms and their impact on certain evolutionary analyses, for consideration of the readers and users in different research areas.

Age Variation in Source Chronograms

Conflict in estimated ages among alternative studies is common in the literature. See, 398 for example, the robust ongoing debate about crown group age of angiosperms 390 (Barba-Montoya, Reis, Schneider, Donoghue, & Yang, 2018; Magallón, Gómez-Acevedo, 400 Sánchez-Reves, & Hernández-Hernández, 2015; Ramshaw et al., 1972; Sanderson & Dovle, 401 2001; Sauquet, Ramírez-Barahona, & Magallón, 2021). Alternative source chronograms 402 available for the same taxa have potentially been estimated implementing different types of 403 calibrations, which affects the resulting node age estimates. For example, in the DateLife 404 analysis of the Fringillidae shown above, the chronograms from one study (Burns et al., 2014) 405 were inferred using molecular substitution rate estimates across birds (Weir & Schluter, 2008), and have much older age estimates for the same nodes than chronograms that were inferred using fossil calibrations (Fig. 4; Supplementary Figs. S4c, S10). Another source of conflict in estimated node ages can arise from different placements for the same calibration, which would imply fundamentally distinct evolutionary hypotheses (Antonelli et al., 2017). 410 For example, two independent researchers working on the same clade should both carefully 411

select and justify their choices of fossil calibration placement. Yet, if one researcher concludes 412 that a fossil should calibrate the ingroup of a clade, while another researcher concludes that 413 the same fossil should calibrate the outgroup of the clade, the resulting age estimates will 414 differ, as the placement of calibrations as stem or crown group is known to significantly affect 415 estimates of time of lineage divergence (Sauquet, 2013). Finally, placement of calibrations 416 also affects uncertainty of node age estimates. For example, nodes that are sandwiched 417 between a calibrated node and a calibrated root have less freedom of movement and hence 418 narrower confidence intervals (Vos & Mooers, 2004), which inflates precision for nodes 419 without calibrations but does not necessarily improve accuracy of the estimated ages. 420

DateLife's summary chronograms are intended to represent all variation in estimated node ages from source chronograms. Node age distribution ranges allow to visually explore ages from source chronograms individually and contextualize and compare them against other chronograms. Researchers that wish to use summary chronograms in downstream evolutionary analysis may select multiple trees sampled from the summary distribution of node ages, to account for variation in source chronograms.

Primary vs Secondary Calibrations

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DateLife constructs summary chronograms using node ages extracted from existing 428 chronograms, i.e. secondary calibrations. In general, the scientific community has more 429 confidence in chronograms using primary calibrations, where the dated tree is generated from 430 a single analysis where carefully chosen fossil calibrations are the source of absolute time 431 information, than in analyses dated using secondary calibrations (Antonelli et al., 2017; Garzón-Orduña, Silva-Brandão, Willmott, Freitas, & Brower, 2015; Graur & Martin, 2004; 433 Sauguet, 2013; Sauguet et al., 2012; Schenk, 2016; Shaul & Graur, 2002). However, implementation of primary calibrations is difficult: it requires specialized expertise and 435 training to discover, place and apply calibrations appropriately (Hipsley & Müller, 2014; 436 Ksepka et al., 2011). One approach is to use fossils that have been widely discussed and 437

previously curated as calibrations to date other trees (Ksepka et al., 2011; Sauquet, 2013), 438 and making sure that all data reflect a coherent evolutionary history (Sauquet, 2013), as for 439 example done by Antonelli et al. (2017). The Fossil Calibration Database provides data for 440 220 primary calibration points encompassing flowering plants and metazoans, that have been 441 curated by experts and used for dating analysis in peer-reviewed publications (Ksepka et al., 442 2015). This database facilitates the use of expert primary fossil calibrations in new phylogenetic dating analyses. Yet, users still require the expertise to locate and calibrate 444 appropriate nodes in their phylogenies which correspond with fossils available in the database. 446

Recently, Powell, Waskin, and Battistuzzi (2020) showed in a simulation study that 447 secondary calibrations using node ages based on previous molecular clock analyses can be as 448 good as primary calibrations. Using several secondary calibrations (as opposed to just one) 449 can provide sufficient information to alleviate or even neutralize potential biases (Graur & 450 Martin, 2004; Sauquet, 2013; Shaul & Graur, 2002). Our cross validation analysis also 451 provides insight into the application of secondary calibrations. Node ages summarized with 452 DateLife and those from the original studies are well correlated (Supplementary Figs. 453 S5-S23). We also note that DateLife estimates for nodes closer to the root tend to be slightly 454 younger than ages from the original studies. In contrast, nodes closer to the tips tend to be 455 slightly older when estimated using our secondary calibrations than ages from the original 456 studies. The only exception to this trend was observed in Burns et al. (2014) chronogram, 457 which generally displays much younger node ages when estimated using secondary 458 calibrations than the original study (Supplementary Figs. S4c, S10), supporting previous observations (Sauquet et al., 2012; Schenk, 2016). However, these younger dates are more likely an example of how multiple secondary calibrations can correct erroneous estimates, as dates on the Burns et al. (2014) tree were obtained using a single secondary calibration based on a previously estimated molecular evolution rate across birds from Weir and 463 Schluter (2008), and appear as major outliers compared to alternate estimates for the same

nodes based on primary fossil calibrations (Fig. 3, Supplementary Fig. S3).

Further research is needed to fully understand the effects of using secondary
calibrations and the use of resulting chronograms in downstream analyses (Hipsley & Müller,
2014; Powell et al., 2020; Schenk, 2016; Shaul & Graur, 2002).

Sumarizing Chronograms

By default, DateLife currently summarizes all source chronograms that overlap with at
least two species names. Users can exclude source chronograms if they have reasons to do so.
Strictly speaking, a good chronogram should reflect the real time of lineage divergence
accurately and precisely. To our knowledge, there are no tested measures to determine
independently when a chronogram is better than another. Yet, several characteristics of the
data used for dating analyses, as well as from the output chronogram itself, could be used to
score the quality of source chronograms.

Some measures that have been proposed are the proportion of lineage sampling and the number of calibrations used (Magallón, 2010; Magallón et al., 2015). 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 fossils used as calibrations, support for nodes and ages, and magnitude of confidence intervals.

DateLife provides an opportunity to capture concordance and conflict among date
estimates, which can also be used as a metric for chronogram reliability. Its open database of
chronograms allows other researchers to do such analyses themselves reproducibly, and
without needing permission. Though, of course, they should follow proper citation practices,
especially for the source chronogram studies.

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The exercise of summarizing age data from across multiple studies provides the 489 opportunity to work with a chronogram that reflects a unified evolutionary history for a 490 lineage, by putting together evidence from different hypotheses. The largest, and 491 taxonomically broadest chronogram currently available from OpenTree was constructed 492 summarizing age data from 2,274 published chronograms using NCBI's taxonomic tree as 493 backbone (Hedges et al., 2015). A summarizing exercise may also amplify the effect of 494 uncertainty and errors in source data, and blur parts of the evolutionary history of a lineage 495 that might only be reflected in source chronograms and lost on the summary chronogram 496 (Sauguet et al., 2021). 497

Effects of Taxon Sampling on Downstream Analyses

Analysis of species diversification of simulated and empirical phylogenies suggest that 499 using a more completely sampled phylogeny provides estimates that are closer to the true 500 diversification history than when analysing incompletely sampled phylogenies (Chang, 501 Rabosky, & Alfaro, 2020; Cusimano, Stadler, & Renner, 2012; Sun et al., 2020). Ideally, 502 phylogenies should be completed using genetic data, but this is a time-consuming and difficult task to achieve for many biological groups. Hence, DateLife's workflow features different ways of assigning divergence times to taxa with missing the absence of branch length data and calibrations and branch lengths for certain taxa.

Completing a phylogeny using a stochastic birth-death polytomy resolver and a 507 backbone taxonomy is a common practice in scientific publications: Jetz et al. (2012), 508 created a chronogram of all 9.993 bird species, where 67% had molecular data and the rest was simulated; Rabosky et al. (2018) created a chronogram of 31,536 ray-finned fishes, of 510 which only 37% had molecular data; Smith and Brown (2018) constructed a chronogram of 511 353,185 seed plants where only 23% had molecular data. These stochastically resolved 512 chronograms return diversification rates estimates that appear less biased than those 513 estimated from their incompletely sampled counterparts, even with methods that account for 514

missing lineages by using sampling fractions (Chang et al., 2020; Cusimano et al., 2012), but can also introduce spurious patterns of early bursts of diversification (Cusimano & Renner, 2010; Sun et al., 2020).

Taxonomy-based stochastic polytomy resolvers also introduce topological differences in 518 phylogenetic trees. The study of macroevolutionary processes largely depends on an 519 understanding of the timing of species diversification events, and different phylogenetic and 520 chronogram hypothesis can provide very different overviews of the macroevolutionary history 521 of a biological group. For example, alternative topologies in chronograms from the same 522 biological group can infer very different species diversification patterns (Rabosky, 2015; Title 523 & Rabosky, 2016). Similarly, there are worries that patterns of morphological evolution 524 cannot be accurately inferred with phylogenies that have been resolved stochastically over a 525 taxonomic backbone, as any patterns would be erased by randomization (Rabosky, 2015). 526 We note that the same applies for geography- and morphology-dependent diversification 527 analysis. Hence, we suggest that phylogenies that have been processed with taxonomy-based 528 stochastic polytomy resolvers, including certain summary chronograms from a DateLife analysis, can be useful as null or neutral models, representing the case of a diversification process that is independent of traits and geographical scenario. 531

Taxonomy-based stochastic polytomy resolvers have been used to advance research in
evolution, still, risks come with this practice. Taken to the extreme, one could generate a
fully resolved, calibrated tree of all modern and extinct taxa using a single taxonomy, a
single calibration, and assigning branch lengths following a birth-death diversification model.
Clearly, this can lead to a misrepresentation of the true evolutionary history. We urge
DateLife users to follow the example of the large tree papers cited above, by carefully
considering the statistical assumptions being made, potential biases, and assessing the
consistency of DateLife's results with prior work.

Knowledge of the evolutionary time frame of organisms is key to many research areas:
trait evolution, species diversification, biogeography, macroecology and more. It is also
crucial for education, science communication and policy, 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 skills to construct
them on their own. Importantly, years of primarily publicly funded research have resulted in
vast amounts of chronograms that are already available in scientific publications, but
functionally hidden from the public and scientific community for reuse.

The DateLife project allows for easy and fast summarization of public and 549 state-of-the-art data on time of lineage divergence. It is available as an R package, and as a 550 web-based R shiny application at www.datelife.org. DateLife provides a straightforward way 551 to get an informed picture of the state of knowledge of the time frame of evolution of 552 different regions of the tree of life, and allows identifying regions that require more research, 553 or that have conflicting information. Additionally, both summary and newly generated trees 554 using the DateLife workflow are useful to evaluate evolutionary hypotheses in different areas of research. We hope that the DateLife project will increase awareness of the existing variation in expert estimations of time of divergence, and foster exploration of the effect of 557 alternative divergence time hypotheses on the results of analyses, nurturing a culture of more cautious interpretation of evolutionary results. 559

Availability

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The DateLife software is free and open source. It can be used online through its R
shiny web application hosted at http://www.datelife.org, and locally through the datelife
R package, available from Zenodo (https://doi.org/10.5281/zenodo.593938 and the CRAN
repository (Sanchez-Reyes et al., 2022). DateLife's web application is maintained using
RStudio's shiny server and the shiny package open infrastructure, as well as Docker and
OpenTree's infrastructure (datelife.opentreeoflife.org). datelife's stable version can be

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installed from the CRAN repository using the command install.packages(pkgs = "datelife") from within R. Development versions are available from DateLife's GitHub repository (https://github.com/phylotastic/datelife) and can be installed using the command devtools::install github("phylotastic/datelife").

Supplementary Material

Supplementary Figures can be viewed and downloaded from their Zenodo repository 572 (https://doi.org/10.5281/zenodo.6683667). Supplementary material, including code, 573 biological examples, benchmark results, data files and online-only appendices, can be 574 downloaded from the Dryad data repository (https://doi.org/10.5061/dryad.cnp5hqc6w), as 575 well as in the Zenodo stable repositories that host the reproducible manuscript 576 (https://doi.org/10.5281/zenodo.7435094), the biological examples 577 (https://doi.org/10.5281/zenodo.7435101), and the software benchmark 578 (https://doi.org/10.5281/zenodo.7435106). Development versions corresponding to all of the above are hosted on GitHub, accesible at https://github.com/LunaSare/datelifeMS1, https://github.com/LunaSare/datelife examples, and https://github.com/LunaSare/datelife benchmark.

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