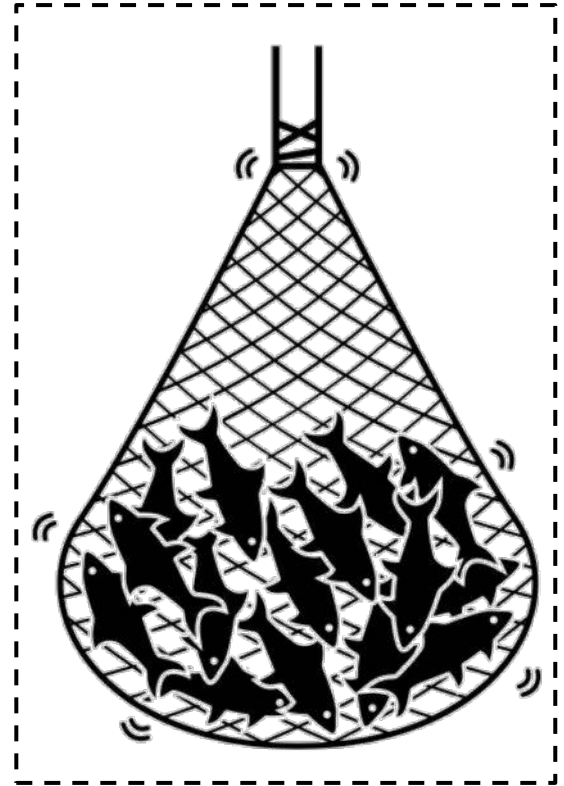


Target Capture Sequencing Approaches

TCS Approaches in Phylogenomics



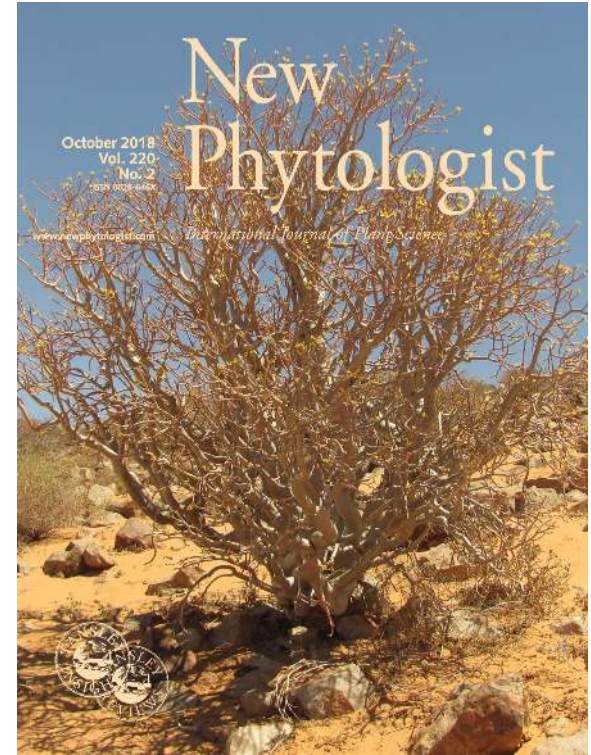
An aerial photograph of the University of Cambridge campus. The image captures a dense cluster of historic stone buildings, many with red-tiled roofs, interspersed with lush green lawns and mature trees. Some trees show early autumn colors. A prominent feature is the tall, white spire of King's College Chapel, which stands out against the sky. In the foreground, a road with a few cars and a bus is visible, along with a large, open green area. The overall scene depicts a well-maintained and historic academic environment.

[illegible]

I got my PhD in Biology @ **Duke University** (USA).
My postdoctoral trajectory spans Madrid's Royal
Botanic Garden (**RJB-CSIC**, ES)



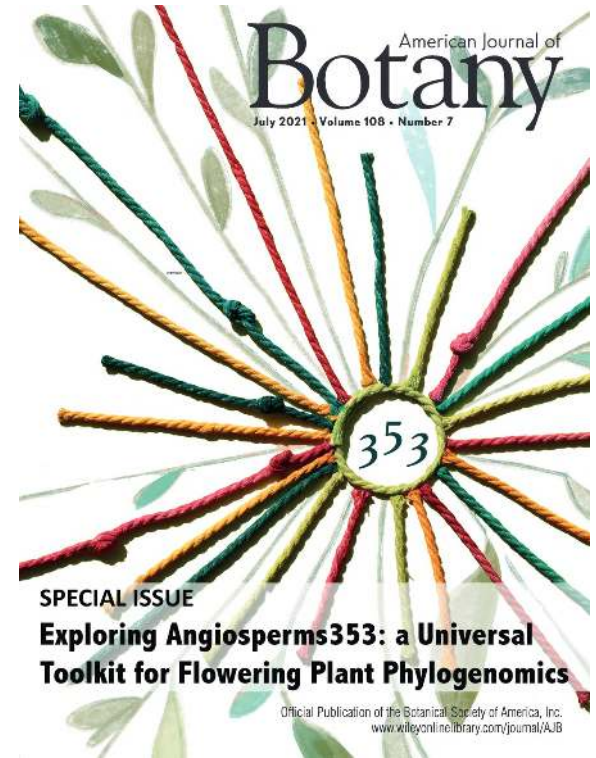
—**Lisa Pokorny, PhD**



I got my PhD in Biology @ **Duke University** (USA).
My postdoctoral trajectory spans Madrid's Royal Botanic Garden (**RJB-CSIC**, ES), Royal Botanic Gardens, Kew (**RBGK**, UK)



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—**Lisa Pokorny, PhD**



I got my PhD in Biology @ **Duke University** (USA).

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Nowadays, I am a **Ramón y Cajal Research Fellow** working for the Spanish National Research Council (CSIC) and based at **RJB** (Madrid, ES).

—**Lisa Pokorny, PhD**

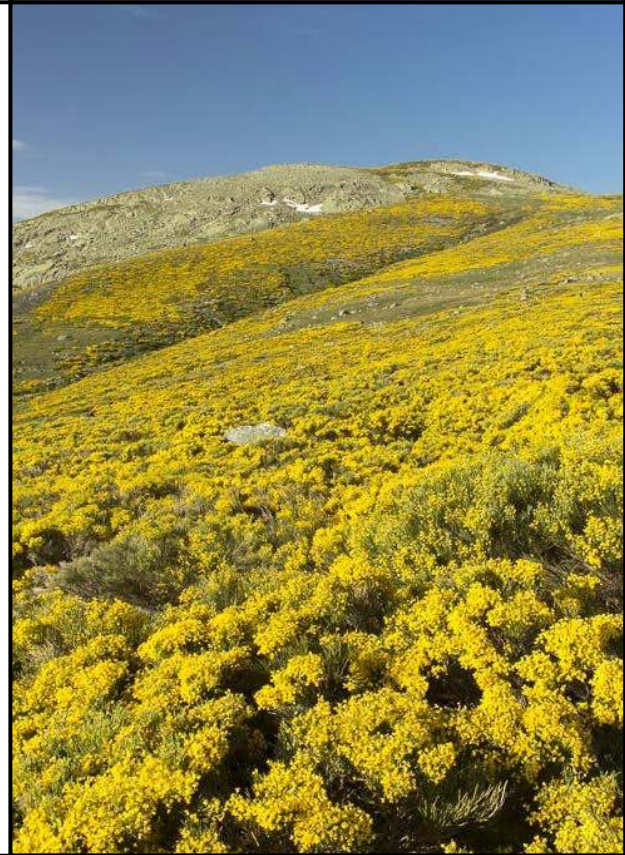


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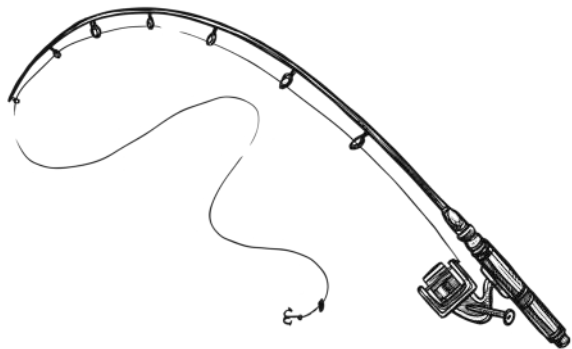
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Biodiversity Genomics

Natural History Collections

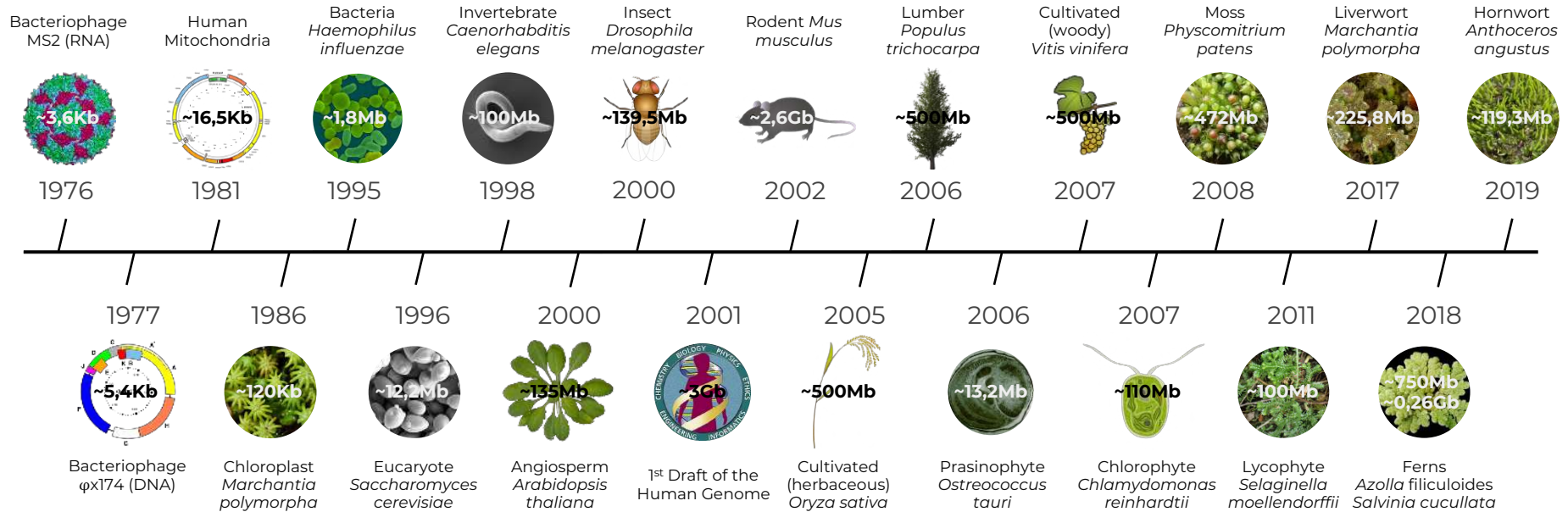


Genome Subsampling

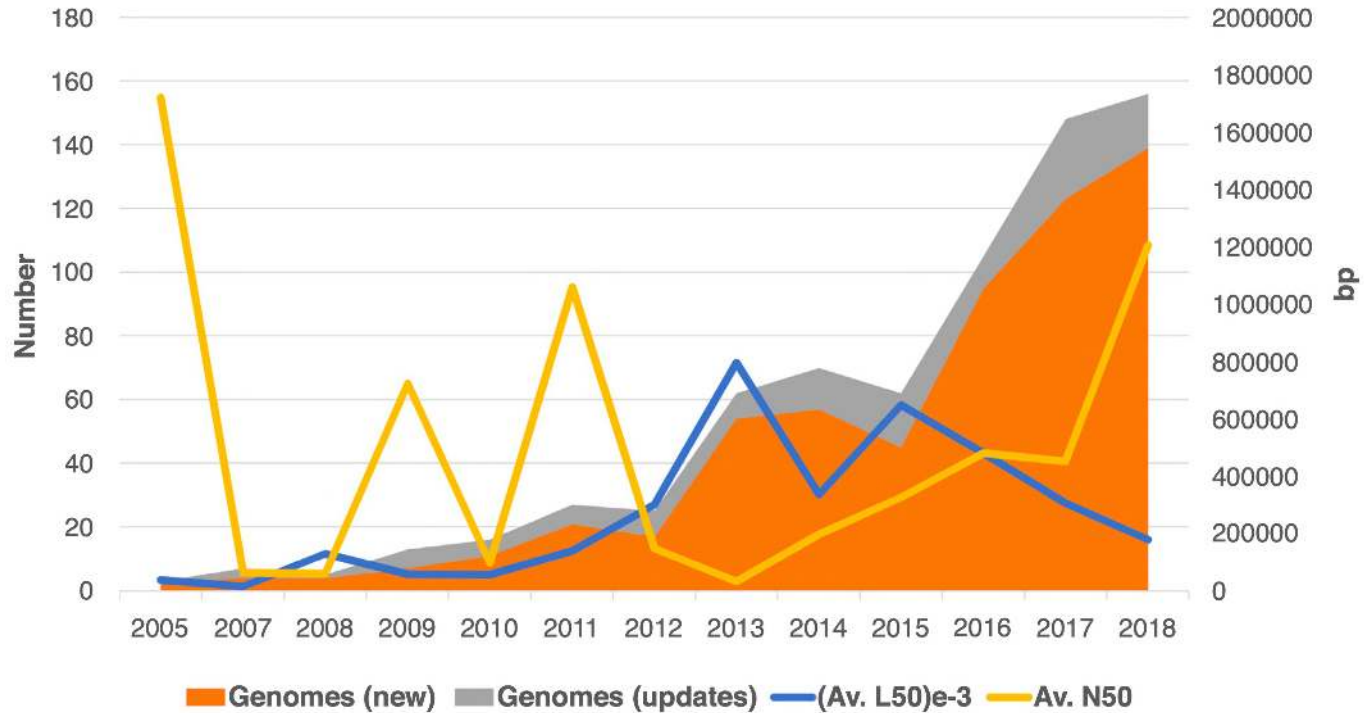
An Introduction to TCS Approaches

01

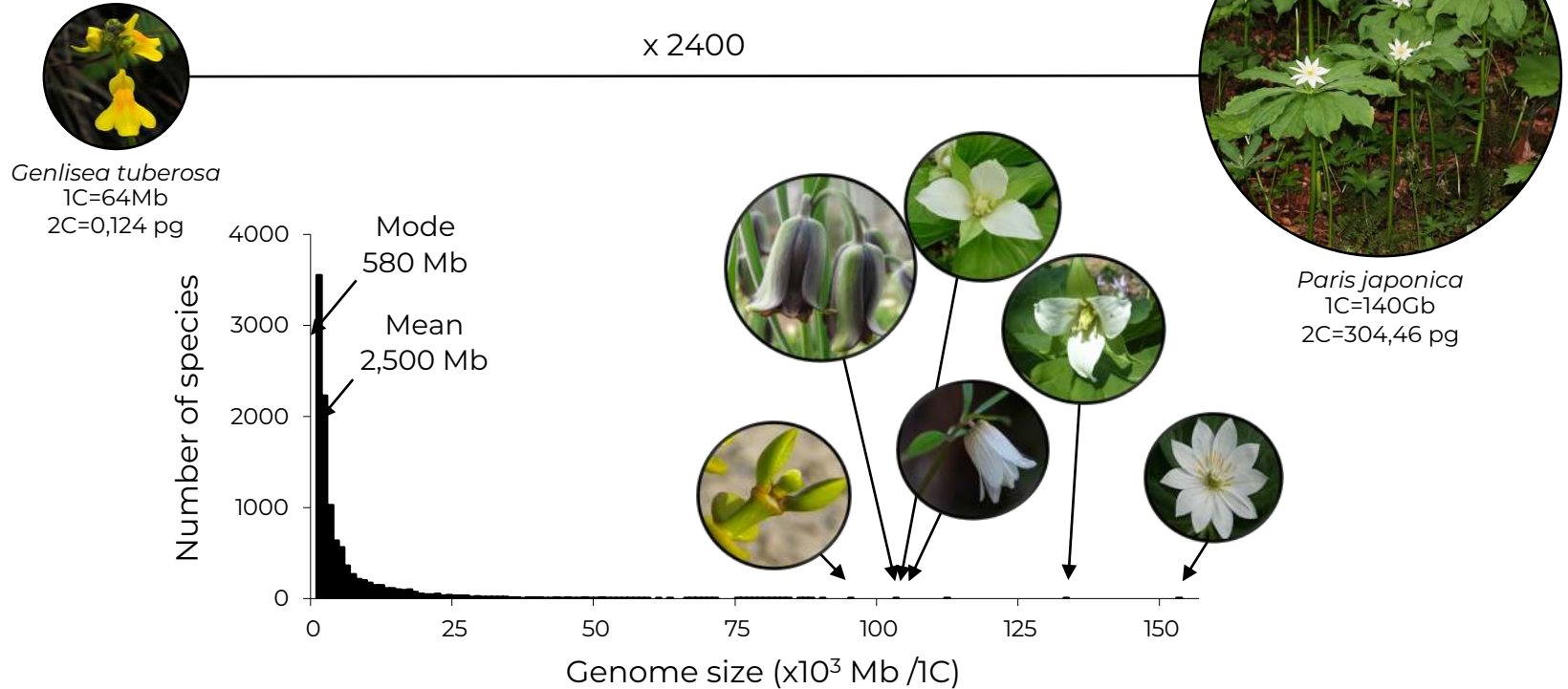
Whole Genome Sequencing (WGS)



Pace of WGS in Plants



Genome Size



Seq Platforms & Costs

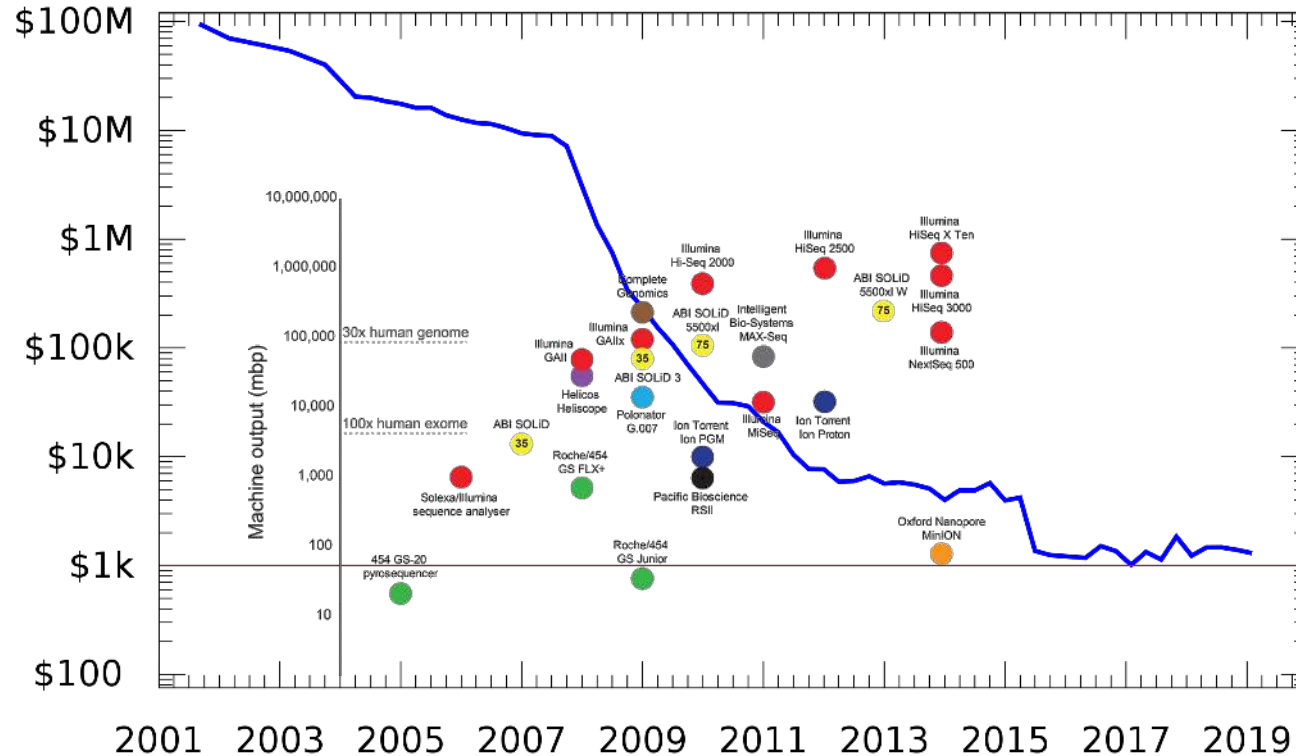
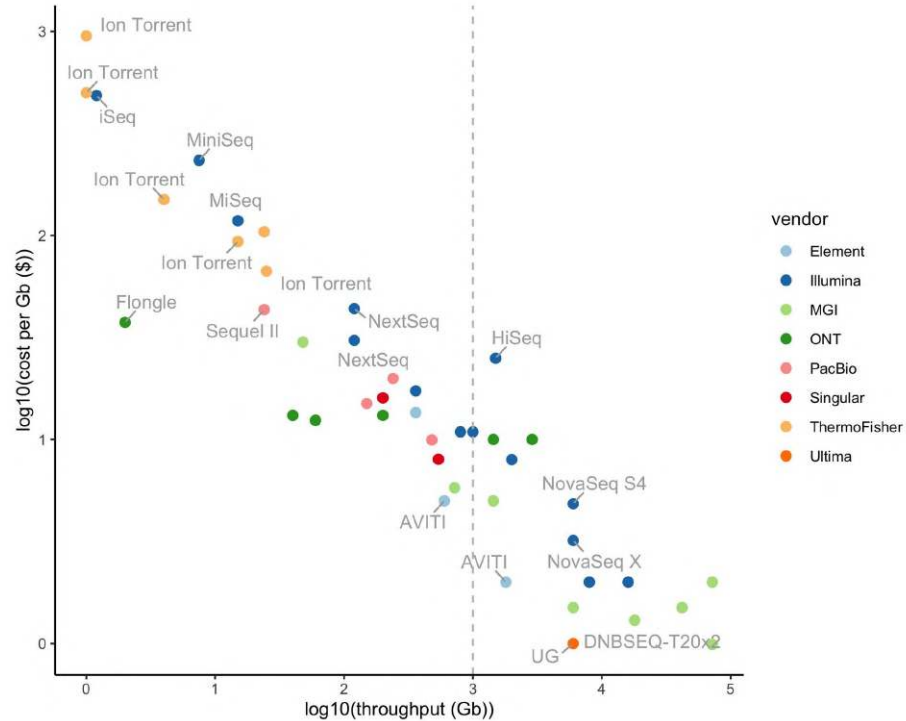


Fig. 1. Reuter et al. 2015. Mol. Cell 58: 586–597 & genome.gov/sequencingcosts

Seq Platforms & Performance

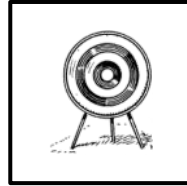


Genome Subsampling



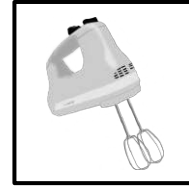
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Genome Skimming
RRS (RAD-seq, GBS)



Targeted

RNA-seq
Target Capture Seq
(TCS)



Mixed

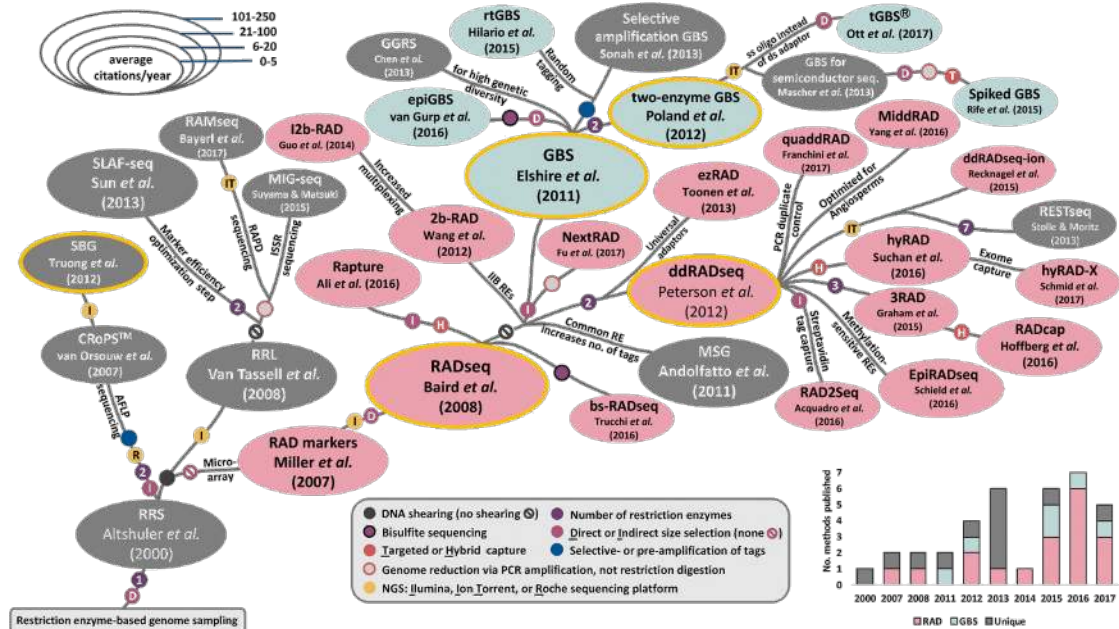
Hyb-Seq

Genome Subsampling



Anonymous

Genome Skimming
RRS (RAD-seq, GBS)

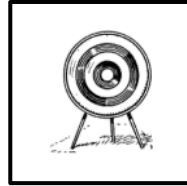


Genome Subsampling



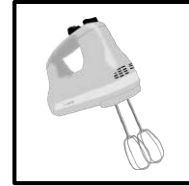
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Genome Skimming
RRS (RAD-seq, GBS)



Targeted

RNA-seq
Target Capture Seq
(TCS)



Mixed

Hyb-Seq

Genome Subsampling

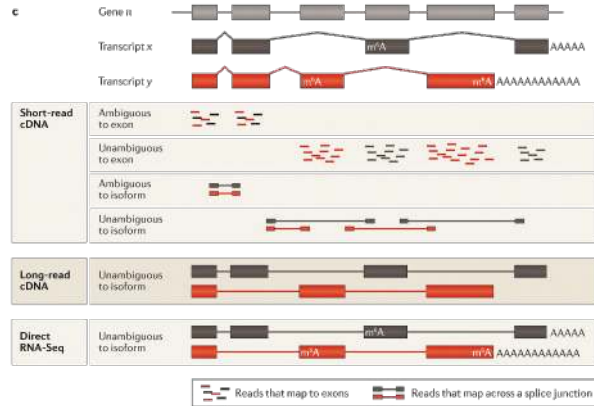
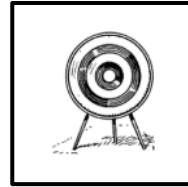


Fig. 1c. Stark et al. 2019. *Nat. Rev. Genet.* 20: 631–656



Targeted RNA-seq Target Capture Seq (TCS)

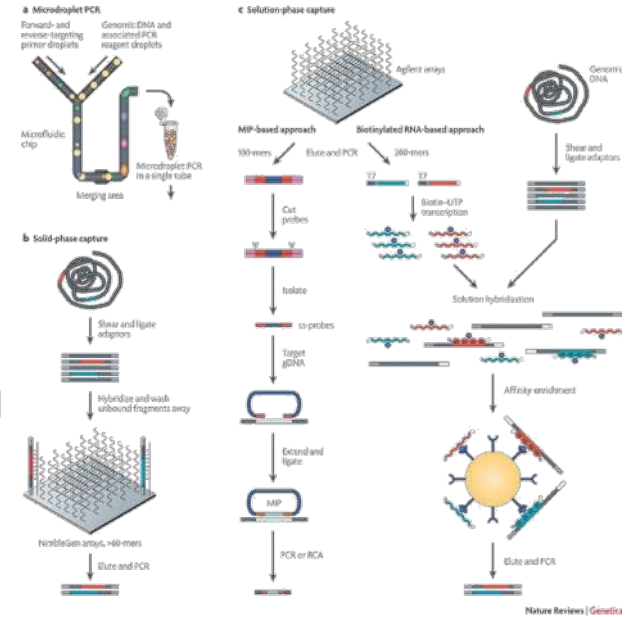


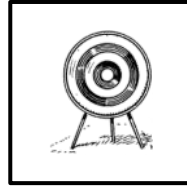
Fig. 5. Metzker. 2010. *Nat. Rev. Genet.* 11: 31–46

Genome Subsampling



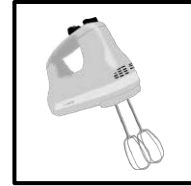
Anonymous

Genome Skimming
RRS (RAD-seq, GBS)



Targeted

RNA-seq
Target Capture Seq
(TCS)



Mixed

Hyb-Seq

Genome Subsampling



Hyb-Seq: Combining Target Enrichment and Genome Skimming for Plant Phylogenomics

Author(s): Kevin Weitemier, Shannon C. K. Straub, Richard C. Cronn, Mark Fishbein, Roswitha Schmickl, Angela McDonnell, and Aaron Liston

Source: Applications in Plant Sciences, 2(9)

Published By: Botanical Society of America

<https://doi.org/10.3732/apps.1400042>

URL: <http://www.bioone.org/doi/full/10.3732/apps.1400042>



Mixed

Hyb-Seq

Genome Subsampling

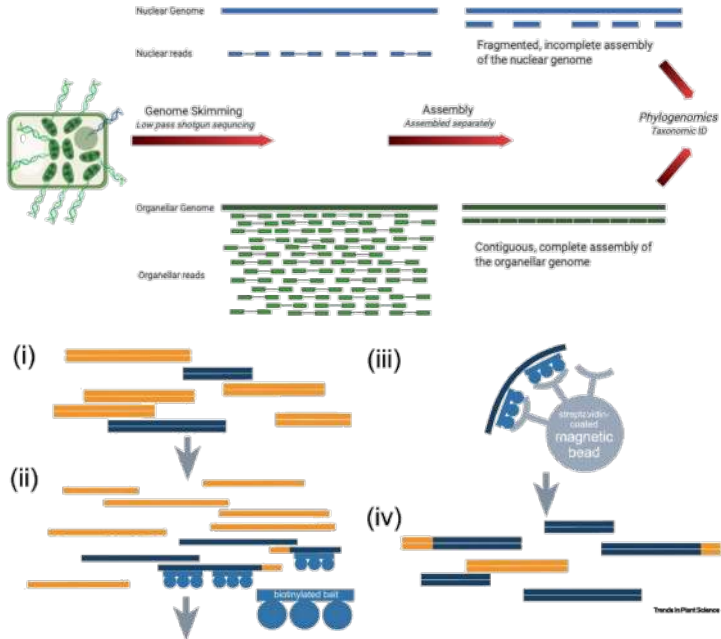


Fig. 1. Dodsworth, Pokorny, Johnson et al. 2019. *Trends Plant Sci.* 24(10): 887–891

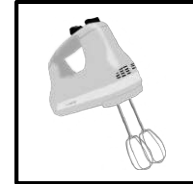
TRPLSC 1851 No. of Pages 4

Trends in Plant Science

Forum

Hyb-Seq for Flowering
Plant Systematics

Steven Dodsworth^{1,2,7,*}
Lisa Pokorny,^{1,3,7}
Matthew G. Johnson,^{4,5,7}
Jan T. Kim,¹ Olivier Maurin,¹
Norman J. Wickett,^{5,6}
Felix Forest,¹ and William J. Baker¹



Mixed
Hyb-Seq

Hyb-Seq Workflow

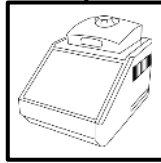
DNA extraction

Comercial kits vs.
in-house protocols



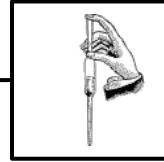
Hybridization

Target enrichment with
probe set



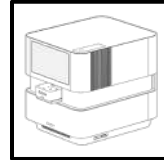
Genomic Library Preparation

Dual indexing for
multiplexing



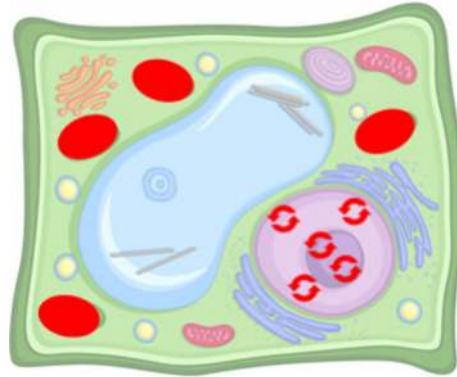
HTS

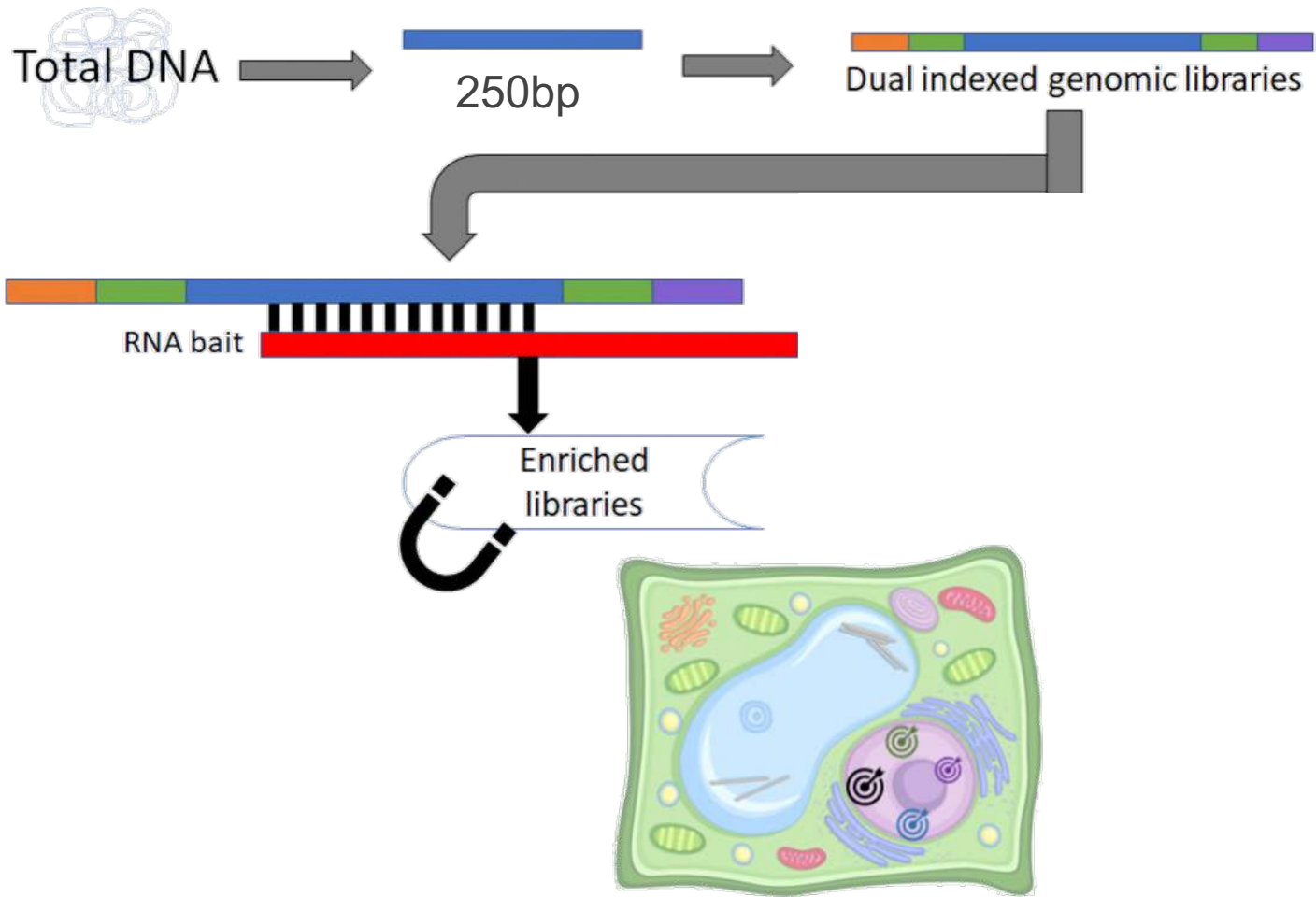
Multiplexed sequencing

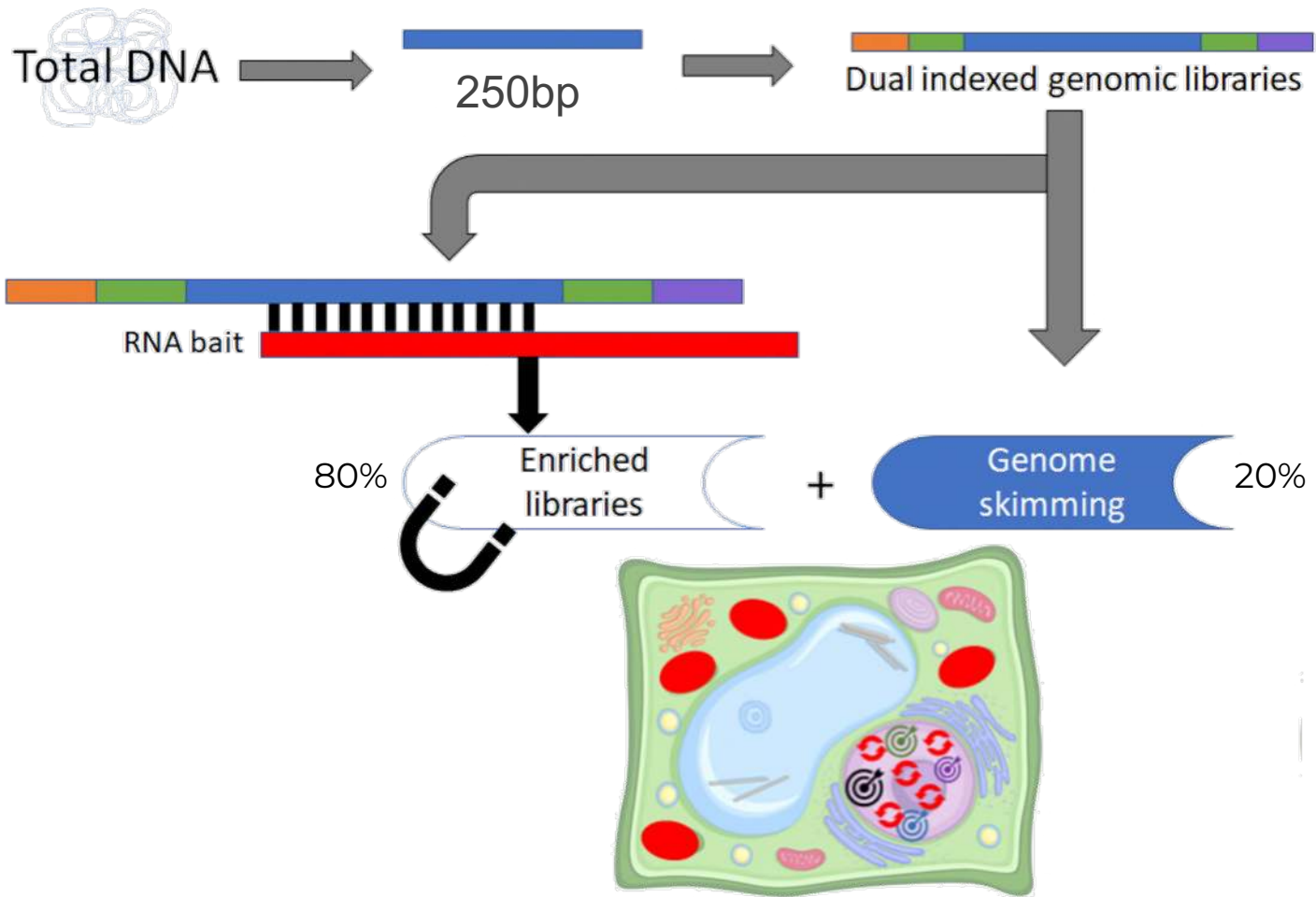




Genome
skimming







Advantages & Disadvantages

Phylogenomics approach	Genomic resources required	Initial bioinformatic investment	Ultimate bioinformatic investment	Initial laboratory cost	Ultimate cost per sample	Low-copy nuclear genes retrieved
<i>Genome skimming</i>	No	None	Medium	Low	Medium	No/Limited
<i>RAD-Seq</i>	No, but helpful	Medium	High	High	Low	No/SNPs
<i>RNA-Seq</i>	No, but helpful	Low	High	Low	High	Yes-thousands
<i>Hyb-Seq</i>	Varies ^b	High ^b	Medium	Low ^b	Medium	Yes-variable

^aInitial costs include the one-time or limited purchase of expensive consumables (e.g., biotinylated baits or adapter sequences). Boxes are highlighted from unfavourable (red) to favourable (green) under each column.

^bIf designing new kit(s) genome or transcriptome resources are required, otherwise readily available kits exist for different groups of plants as well as angiosperms as a whole (Angiosperms-353) and are much cheaper than designing a new custom bait set.

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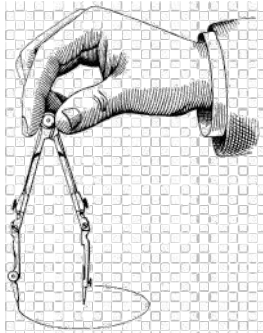
Implementing TCS

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Biodiversity Genomics

Natural History Collections

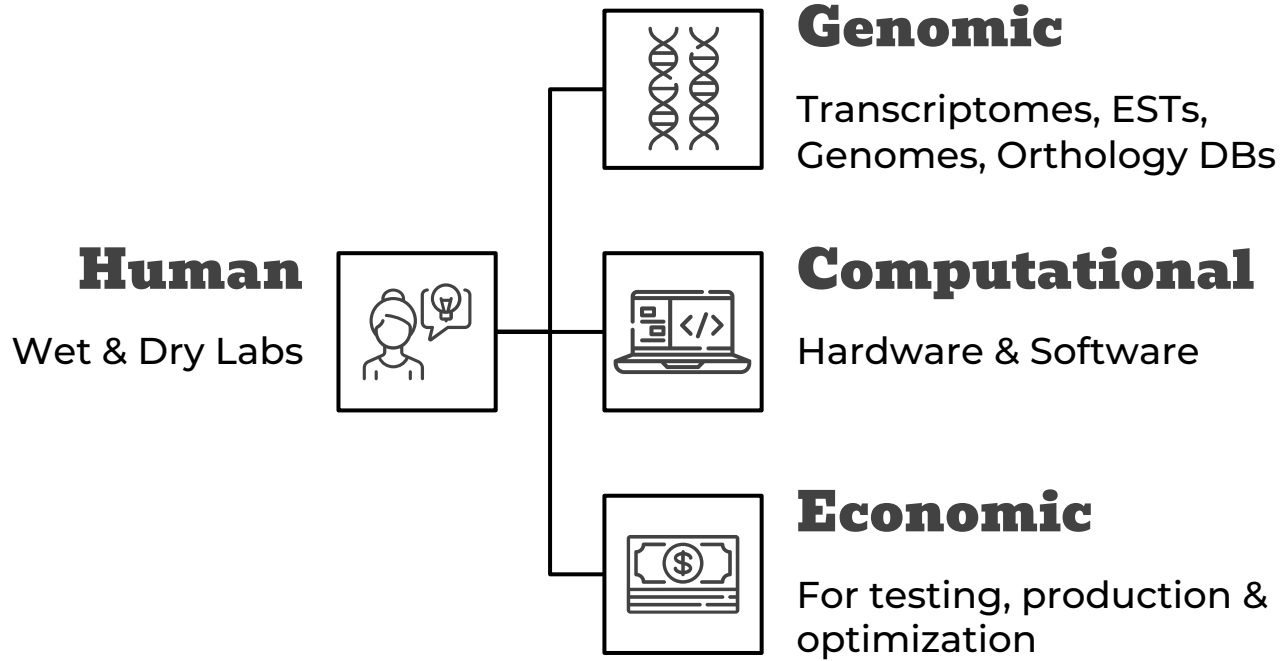


Probe Set Design

Custom Kits

02

Resources needed



Resources needed

Human

Wet & Dry Labs

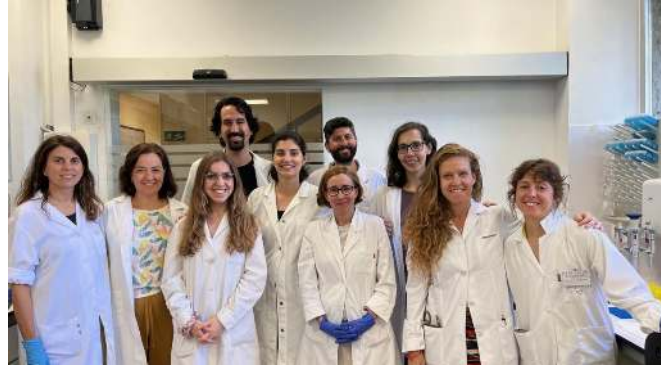


Human Resources



Wet Lab

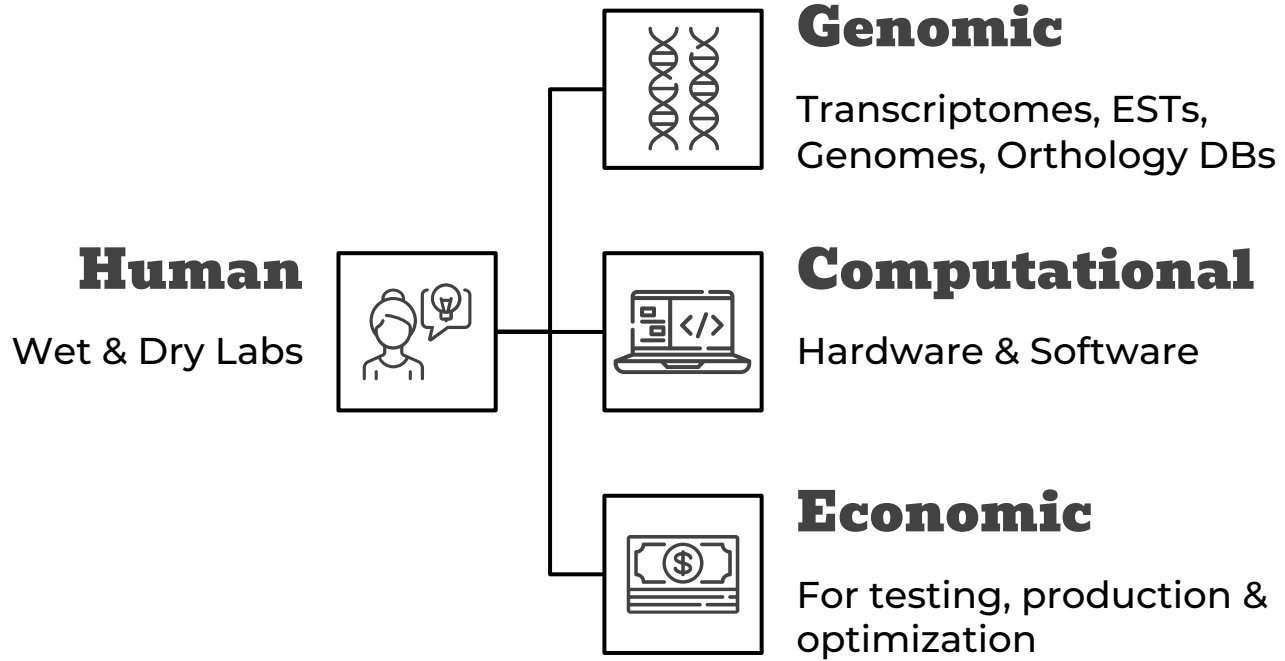
REAL JARDÍN
BOTÁNICO
 **CSIC**
CONSEJO SUPERIOR DE INVESTIGACIONES CIENTÍFICAS



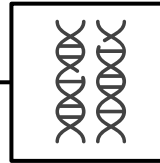
Dry Lab



Resources needed



Resources needed



Genomic

Transcriptomes, ESTs,
Genomes, Orthology DBs

Transcriptomes



Phylotranscriptomic analysis of the origin and early diversification of land plants

Norman J. Wickett^{1,4,12}, Sowash Miralamb¹, Nam Nguyen¹, Tandy Warnow¹, Eric Carpenter¹, Naim Mataszi¹, Saravanan Ayyampalayam¹, Michael S. Barker¹, J. Gordon Burleigh¹, Matthew A. Gitzendanner¹, Brad R. Ruhfel^{1,4}, Eric Wafarua¹, Joshua P. Der¹, Sean W. Graham¹, Sarah Mathew¹, Michael Melkonian¹, Douglas E. Soltis^{1,4}, Pamela S. Soltis^{1,4}, Nicholas W. Miles¹, Carl J. Rothfels^{1,4}, Xia Pokorny¹, A. Jonathan Shaw¹, Lisa DeGironimo¹, Dennis W. Stevenson¹, Barbara Seidel¹, Juan Carlos Villarreal¹, Beatrice Roure¹, Hervé Philippe^{1,2}, Claude W. dePamphilis¹, Tao Chen¹, Michael K. Deyholos¹, Regina S. Baucum¹, Toni M. Kutcher¹, Megan M. Augustin¹, Jun Wang¹, Yong Zhang¹, Zhijian Tian¹, Zhiyong Yang¹, Xiaohu Wu¹, Pao-Sun Gan¹, Garne Ka-Shu Wong^{1,2}, and James Leebens-Mack^{1,2}

Article

One thousand plant transcriptomes and the phylogenomics of green plants

<https://doi.org/10.1038/s41588-019-1693-2>

One Thousand Plant Transcriptomes Initiative

Received: 17 November 2017

Accepted: 12 September 2016

Published online: 23 October 2015

Организмический

Green plants (Viridiplantae) include around 450,000–500,000 species² of great diversity and have important roles in terrestrial and aquatic ecosystems. Here, as part of the One Thousand Plant Transcriptomes Initiative, we sequenced the vegetative transcriptomes of 1,124 species that span the diversity of plants in a broad sense (Archaeplastida), including green plants (Viridiplantae), glaucophytes (Glaucophyta) and red algae (Rhodophyta). Our analysis provides a robust phylogenetic framework for examining the evolution of green plants. Most inferred species relationships are well supported across multiple species tree and supermatrix analyses, but discordance among plastid and nuclear gene trees at a few important nodes highlights the

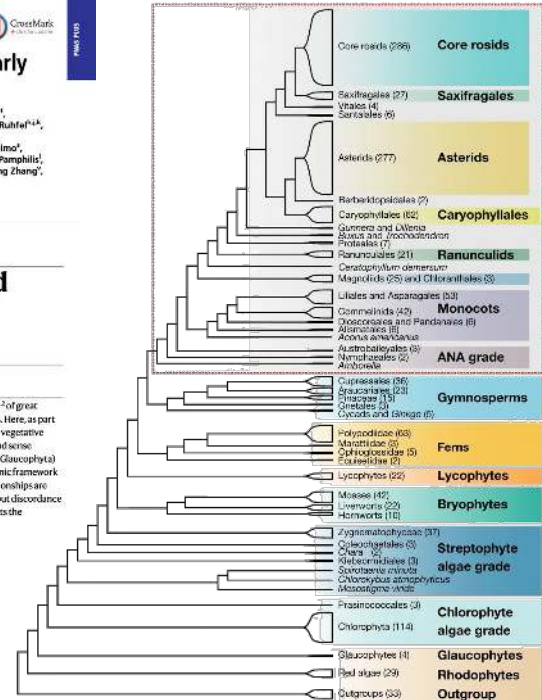
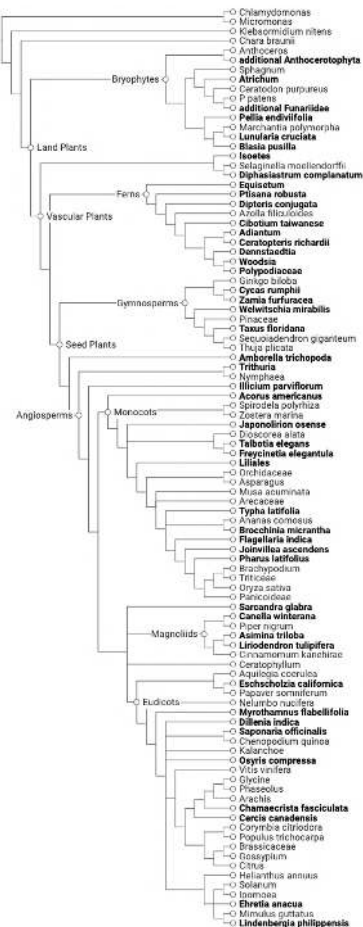


Fig. 1b. OTPTI. 2019. Nature 574: 679–685.

Genomes



OGG

OPEN GREEN GENOMES INITIATIVE

Welcome to **OGG**

Overview

Release Notes

News



The Open Green Genomes Initiative (OGG, a Joint Genome Institute **community science program**) seeks to improve the phylogenetic representation of reference genome sequences across land plants by generating at least thirty-five high-quality genome assemblies and annotations across all major evolutionary lineages in the land plant tree of life (target genomes are labelled in **bold** below). This work will greatly improve comparative analyses of the genes, regulatory networks and metabolic pathways influencing plant growth, responses to environmental stress, and production of valuable plant products. The resulting comparative framework will drive translation of genome science into predictive understanding and inform engineering of efficient production of plant-based biofuels and bioproducts.

Sequence a member of every embryophyte genus and a phylodiverse collection of algae/protists

① Community effort to collect the samples/materials worldwide

9000 embryophytes and
4000 algae/protists



10KP

10 KP online sample submission portal

② Nucleic acid extractions

RNA extraction

HMW DNA extraction

③

Library preparation

recruit linked-reads technologies for library construction

10X Genomics or BGI's LFR platform

④

Sequencing using MGISEQ

⑤

Genome assembly & annotation

⑥

Database & analyses

⑦

Data release and publications



CNCB open access database



Biodiversity data release

Orthology Databases



Ensembl

<http://www.ensembl.org/>



EggNOG

<http://eggnog5.embl.de/>



OrthoDB

<https://www.orthodb.org/>



PLAZA

[bioinformatics.psb.ugent.be/
plaza/](http://bioinformatics.psb.ugent.be/plaza/)



MetaPhOres

orthology.phylomedb.org/



OMA

<https://omabrowser.org/>



KO (KEGG)

www.genome.jp/kegg/ko.html



FungiDB

<https://fungidb.org/>

Orthology Databases

D1468–D1474 *Nucleic Acids Research*, 2022, Vol. 50, Database issue
<https://doi.org/10.1093/nar/gkab1024>

Published online 8 November 2021

PLAZA 5.0: extending the scope and power of comparative and functional genomics in plants

Michiel Van Bel^{1,2}, Francesca Silvestri^{1,2}, Eric M. Weitz³, Lukasz Kreft⁴,
Alexander Botzki⁵, Frederik Coppens^{1,2} and Klaas Vandepoele^{1,2,6,*}

¹Department of Plant Biotechnology and Bioinformatics, Ghent University, Technologiepark 71, 9052 Ghent, Belgium, ²VIB Center for Plant Systems Biology, Technologiepark 71, 9052 Ghent, Belgium, ³Data Sciences Platform, Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA, ⁴Institute of Biochemistry and Biophysics, Polish Academy of Sciences, Pawińskiego 5A 02-106 Warsaw, Poland, ⁵VIB Bioinformatics Core, 9052 Ghent, Belgium and ⁶Bioinformatics Institute Ghent, Ghent University, Technologiepark 71, 9052 Ghent, Belgium

Received September 16, 2021; Revised October 12, 2021; Editorial Decision October 12, 2021; Accepted October 13, 2021



PLAZA

[bioinformatics.psb.ugent.be/
plaza/](https://bioinformatics.psb.ugent.be/plaza/)

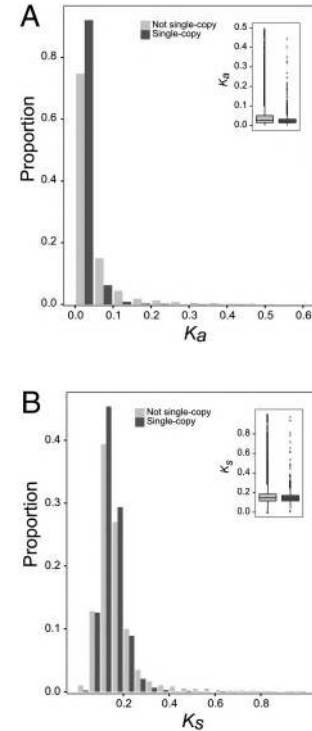
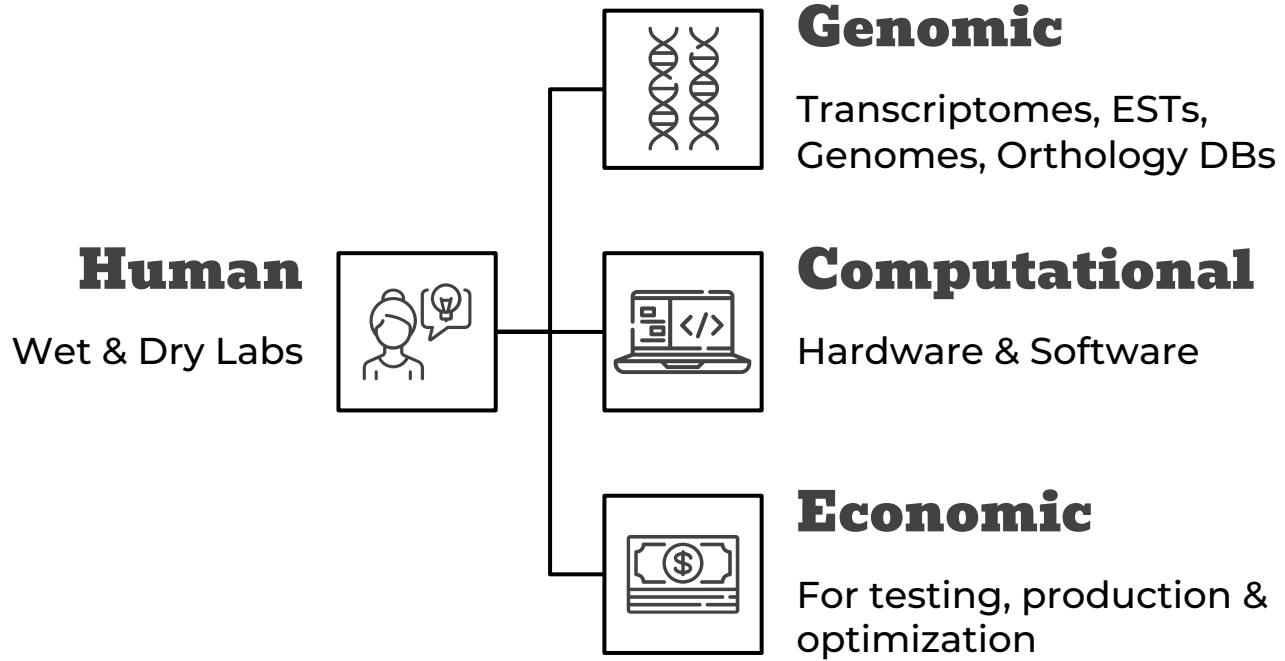
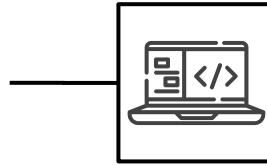


Fig. 3. De Smet et al. 2013. *PNAS* 110: 2898–2903

Resources needed



Resources needed



Computational

Hardware & Software

Hardware



**Barcelona
Supercomputing
Center**

Centro Nacional de Supercomputación



CESGA

GALICIA SUPERCOMPUTING CENTER



Software



Bioinformatics, 2018, 1–4
doi: 10.1093/bioinformatics/bty548
Advance Access Publication Date: 29 June 2018
Applications Note

OXFORD

Software Note | Open Access |

MarkerMiner 1.0: A new application for phylogenetic marker development using angiosperm transcriptomes[†]

Srikanth Chamala , Nicolás García, Grant T. Godden, Vivek Krishnakumar, Ingrid E. Jordon-Thaden, Riet De Smet, W. Brad Barbazuk, Douglas E. Soltis, Pamela S. Soltis

First published: 06 April 2015 | <https://doi.org/10.3732/apps.1400115> | Citations: 32

[†] The authors thank all oneKP contributors, especially Gane Ka-Shu Wong, and BGI. Funding for S.C. was provided by the National Science Foundation (NSF; grant IOS-0922742 [P.S.S., D.E.S., W.B.B.]). Research funding for N.G. and G.T.G. was provided in part by NSF Doctoral Dissertation Improvement grants DEB-1310839 (P.S.S. and N.G.) and DEB-1210671 (P.S.S. and G.T.G.), respectively. The salary of I.J.T. was provided by the David Burpee Endowment and Chris Martine (Bucknell University).

[‡] Authors are listed alphabetically by surname and contributed equally.

BaitFisher: A Software Package for Multispecies Target DNA Enrichment Probe Design

Christoph Mayer, Manuela Sann, Alexander Donath, Martin Meixner, Lars Podsiadlowski, Ralph S. Peters, Malte Petersen, Karen Meusemann, Karsten Lier, Johann-Wolfgang Wägele ... [Show more](#)

[Author Notes](#)

Molecular Biology and Evolution, Volume 33, Issue 7, July 2016, Pages 1875–1886,
<https://doi.org/10.1093/molbev/msw056>

Published: 23 March 2016

Sequence analysis

MrBait: universal identification and design of targeted-enrichment capture probes

Tyler K. Chafin*, Marlis R. Douglas and Michael E. Douglas

Department of Biological Sciences, University of Arkansas, Fayetteville, AR 72701, USA

*To whom correspondence should be addressed.

Associate Editor: John Hancock

Received on April 16, 2018; revised on June 11, 2018; editorial decision on June 27, 2018; accepted on June 28, 2018

Kuchinski et al. *BMC Genomics* (2022) 23:579
<https://doi.org/10.1186/s12864-022-08790-4>

BMC Genomics

RESEARCH

Open Access

ProbeTools: designing hybridization probes for targeted genomic sequencing of diverse and hypervariable viral taxa

Kevin S. Kuchinski^{1,2*}, Jun Duan¹, Chelsea Himsworth^{3,4}, William Hsiao^{1,5} and Natalie A. Prystajek^{1,6}



Software



Software Note | [Open Access](#) | [CC](#) | [BY](#) | [NC](#) | [ND](#)

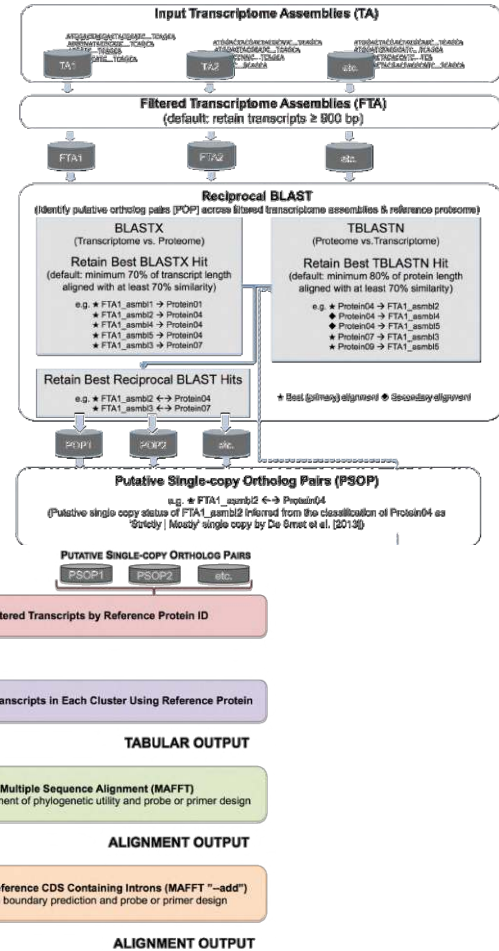
MarkerMiner 1.0: A new application for phylogenetic marker development using angiosperm transcriptomes[†]

Srikanth Chamala , Nicolás García, Grant T. Godden, Vivek Krishnakumar, Ingrid E. Jordon-Thaden, Riet De Smet, W. Brad Barbazuk, Douglas E. Soltis, Pamela S. Soltis

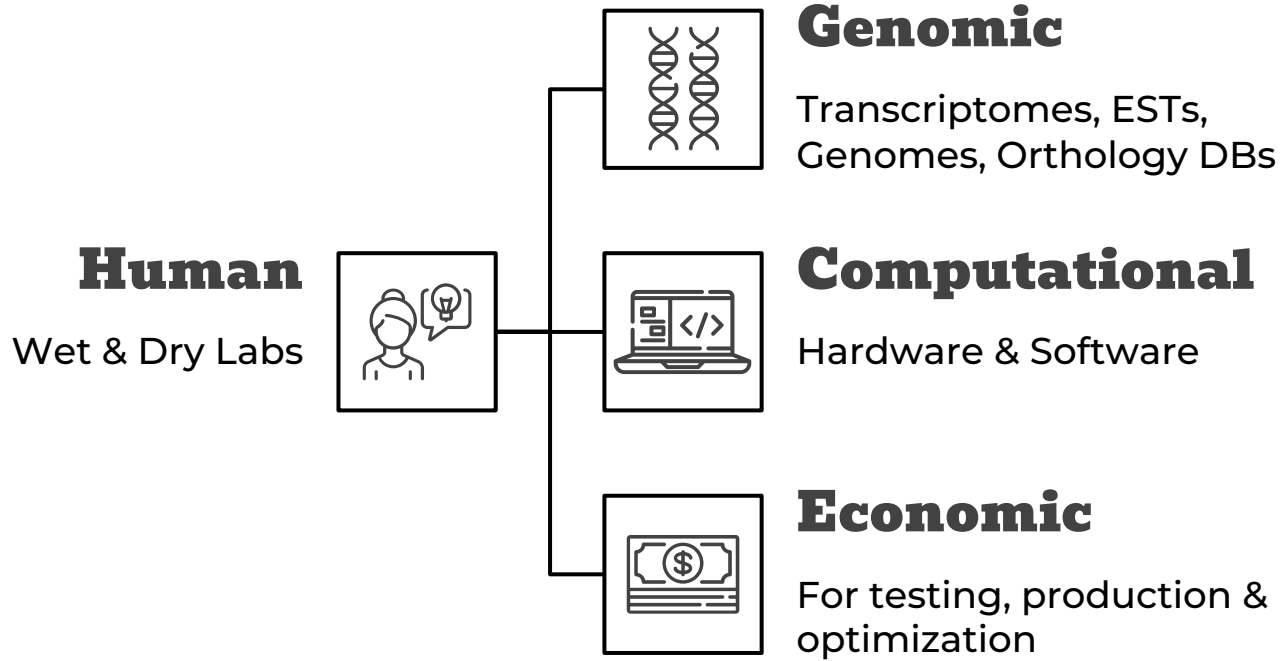
First published: 06 April 2015 | <https://doi.org/10.3732/apps.1400115> | Citations: 32

[†] The authors thank all oneKP contributors, especially Gane Ka-Shu Wong, and BGI. Funding for S.C. was provided by the National Science Foundation (NSF; grant IOS-0922742 [P.S.S., D.E.S., W.B.B.]). Research funding for N.G. and G.T.G. was provided in part by NSF Doctoral Dissertation Improvement grants DEB-1310839 (P.S.S. and N.G.) and DEB-1210671 (P.S.S. and G.T.G.), respectively. The salary of I.J.T. was provided by the David Burpee Endowment and Chris Martine (Bucknell University).

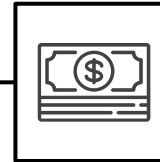
[‡] Authors are listed alphabetically by surname and contributed equally.



Resources needed



Resources needed



Economic

For testing, production & optimization

Economic Resources

TABLE 1. Comparison of consumable costs between conventional and low-cost methods in the target capture workflow. See text for full explanation of methods and alternatives and for equipment cost considerations.^a

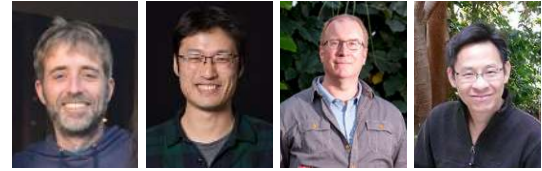
Hyb-Seq procedure	Usual technique	Cost-saving technique	Usual technique price (per sample)	Cost-saving technique price (per sample)	Estimated fold-cost savings
DNA extraction	QIAGEN DNeasy 96	CTAB	\$3.11	\$0.29	10.72
Quality control	TapeStation	Qubit + gel	\$3.21	\$1.30	2.47
Fragmentation	Sonicator	Fragmentase	\$6.76	\$1.41	4.79
Library prep	NEBNext Ultra II DNA full volume	NEB half volume	\$29.20	\$14.60	2.00
Purification	AMPure beads	Homebrew beads	\$2.26	\$0.08	28.25
myBaits	24-plex	24-plex + dilute myBaits	\$2.16	\$0.56	3.86
Sequencing	MiSeq 2 × 300 bp (96-plex)	HiSeq X (384-plex)	\$18.50	\$4.42	4.19
Overall			\$65.20	\$22.66	2.88

^aPrices (in U.S. dollars) are approximate and may vary by institution.

Economic Resources

Duke University

One Thousand Plant Transcriptomes (1KP, Alberta, Musea Ventures, BGI, etc.)



RJB-CSIC

AFFLORA & BAYESNEXT (State Research Agency, AEI)
BOREOTROP (Community of Madrid)



RBGK

Global Tree Seed Bank (GTSB) Programme (Garfield Weston Foundation)
Plant and Fungal Trees of Life (PAFTOL) Programme (Calleva & GW Foundations)



CBGP UPM-INIA/CSIC

Severo Ochoa Excellence Program (AEI)

IBB CSIC-MCNB

GIANTS & CANTHAROMICS (AEI)



Probe Set Design



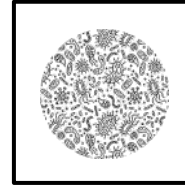
Custom Kits

Euphorbiaceae431

Dioscoreaceae303

Cortinariaceae193

Etc.



Universal Kits

Angiosperms353

ProkaryoKit

Etc.

Euphorbiaceae431

Classification: order Malpighiales
(Eudicot), ~7.5K species (~300 genera)

Importance: rubber (latex), cassava
(edible tuber), castor oil, etc.

Available resources:

2 *Euphorbia* transcriptomes  **1KP**

1 *Ricinus communis* genome



MarkerMiner & Plaza DB

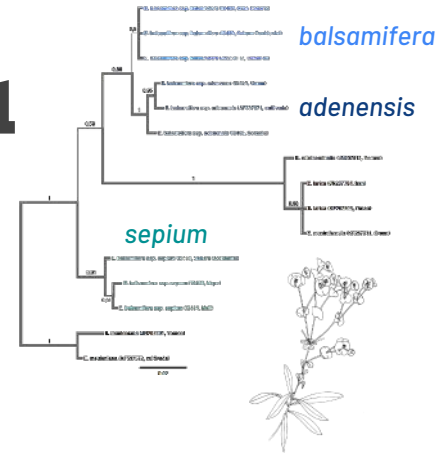
Daicel Arbor Biosciences 

Resulting kit:

431 ~single copy nuclear orthologs

~700 Kbp matrix

~20K 120-mer tiled probes (variable
tiling: T 2x y G 1,5x)



Research

New
Phytologist

Bridging the micro- and macroevolutionary levels in
phylogenomics: Hyb-Seq solves relationships from populations
to species and above

Tamara Villaverde^{1*}, Lisa Pokorný^{2*}, Sanna Olsson³, Mario Rincón-Barrado¹, Matthew G. Johnson^{4,5},
Elliot M. Gardner⁶, Norman J. Wickett^{5,7}, Julià Molero⁸, Ricarda Riina^{1†} and Isabel Sanmartín^{1†}

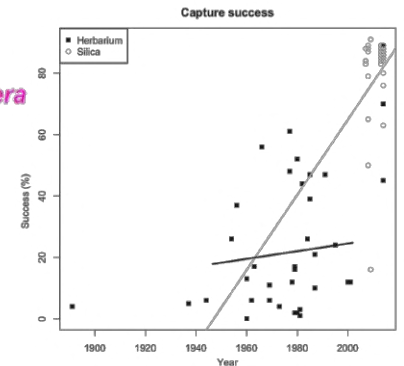
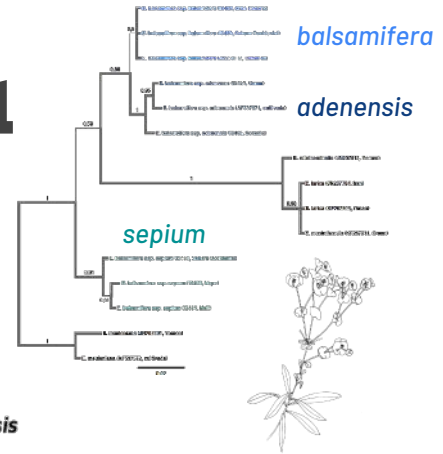
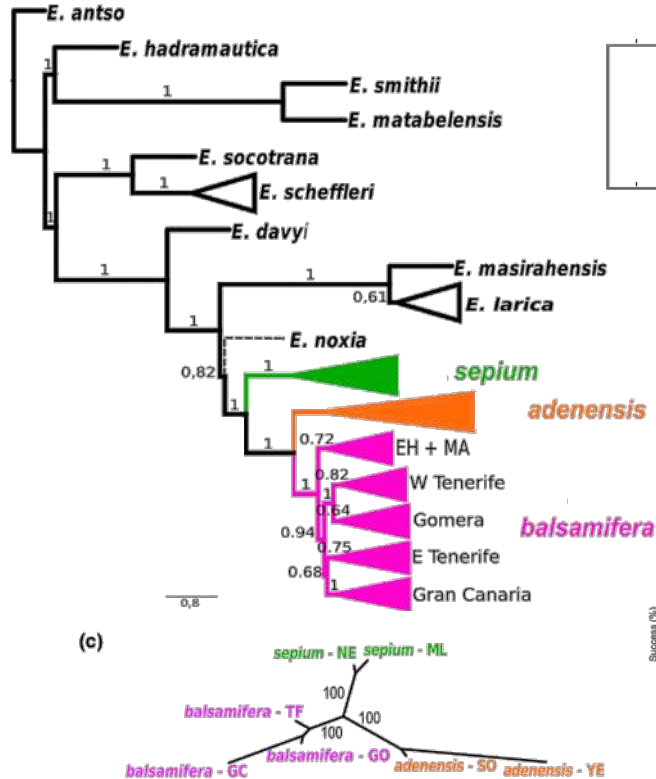
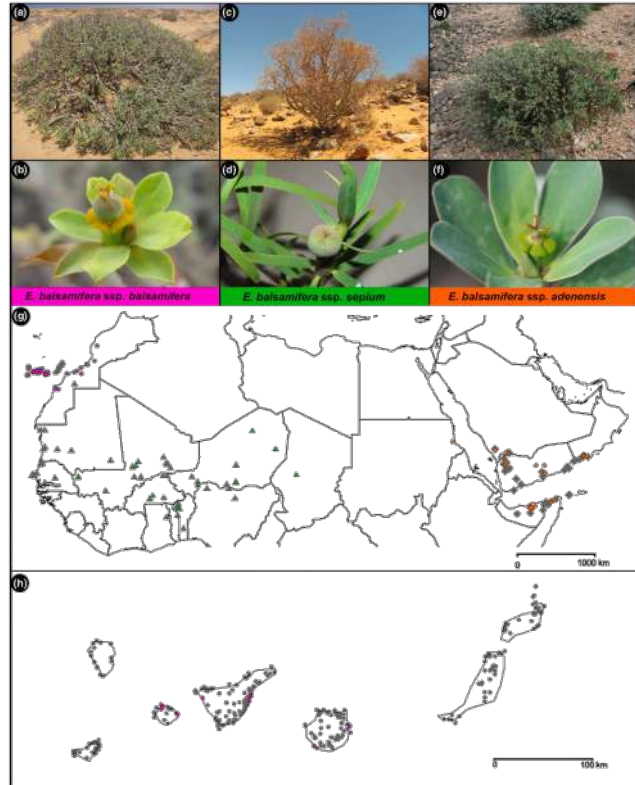
¹Real Jardín Botánico (RJB-CSIC), Plaza de Murillo 2, 28014 Madrid, Spain; ²Comparative Plant and Fungal Biology Department, Royal Botanic Gardens, Kew, Richmond, TW9 3DS, UK;

³Department of Forest Ecology and Genetics, INIA Forest Research Centre (INIA-CIFOR), Ctra. de la Coruña km. 7.5, 28040 Madrid, Spain; ⁴Department of Biological Sciences, Texas Tech University, 2901 Main St, Lubbock, TX 79409-43131, USA; ⁵Department of Plant Science and Conservation, Chicago Botanical Garden, 1000 Lake Cook Road, Glencoe, IL 60022, USA;

⁶The Morton Arboretum, 4100 Illinois 53, Lisle, IL 60532 USA; ⁷Program in Plant Biology and Conservation, Northwestern University, 2205 Tech Drive, Evanston, IL 60208 USA;

⁸Laboratori de Botànica, Departament de Biologia, Sanitat i Medi Ambient, Facultat de Farmàcia, Universitat de Barcelona, 08028 Barcelona, Spain

Euphorbiaceae431



Dioscoreaceae303



Classification: order Dioscoreales
(Monocot), ~300 species (~4 genera)

Importance: yams (edible tuber),
steroidal saponins (contraceptives), etc.

Available resources:  **1KP**

3 *Dioscorea* & 1 *Tacca* transcriptomes

1 *Oryza sativa* & 1 *Xerophyta viscosa*



MarkerMiner
Locus development made easy

& Plaza DB

Daicel Arbor Biosciences 

 Applications
in Plant Sciences



Resulting kit:

260 ~single copy nuclear orthologs

43 genes of agronomic interest

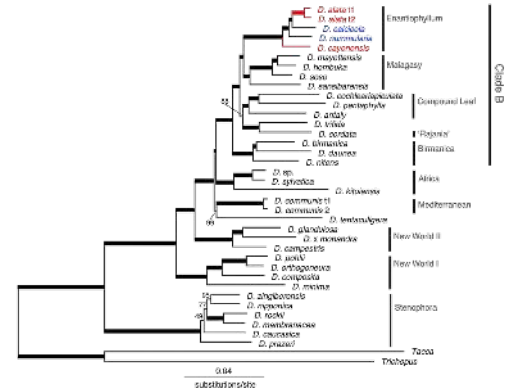
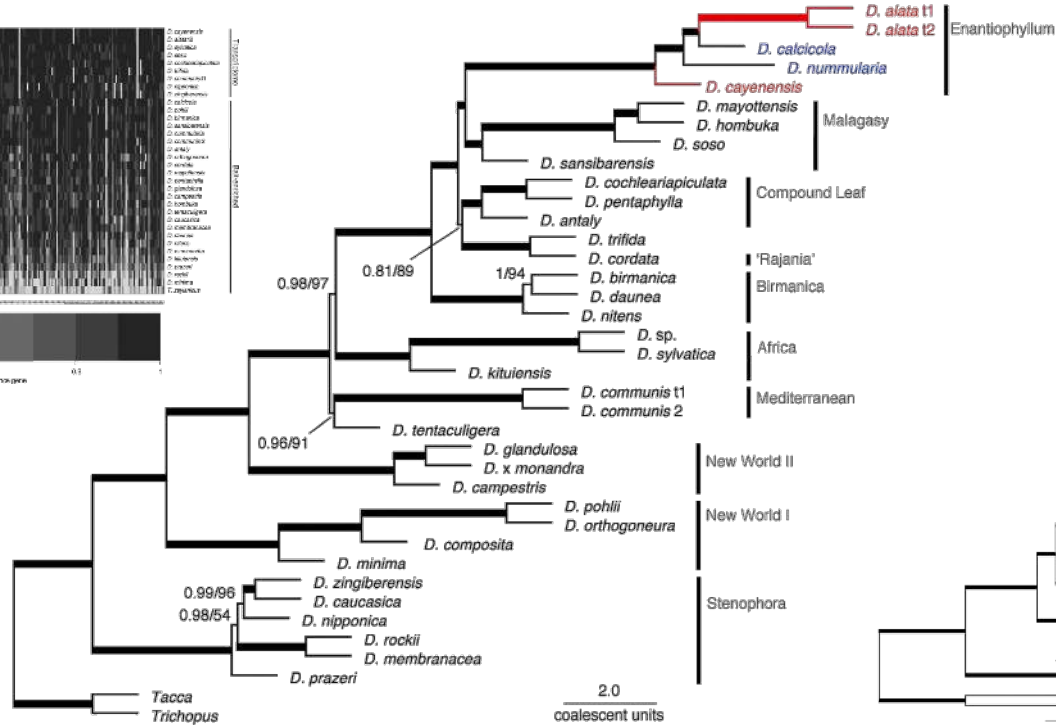
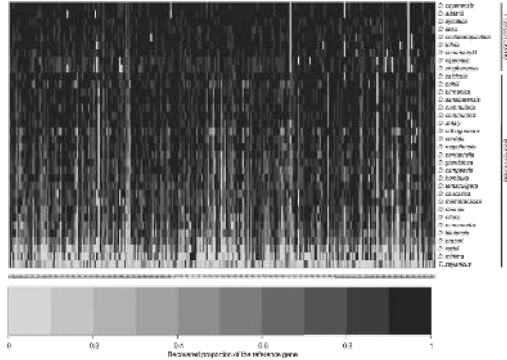
~442 Kbp matrix

~17K 120-mer 3x tiled probes

**A customized nuclear target enrichment approach for
developing a phylogenomic baseline for *Dioscorea* yams
(Dioscoreaceae)**

Marybel Soto Gomez^{1,2,3} , Lisa Pokorny¹ , Michael B. Kantar¹ , Félix Forest¹ , Ilia J. Leitch¹ , Barbara Gravendeel^{1,4,7} ,
Paul Wilkin¹ , Sean W. Graham^{1,2} , and Juan Viruel¹ 

Dioscoreaceae303





Cortinariaceae193

Classification: order Agaricales
(Basidiomycota), ~3K species (1 genus)

Importance: ectomycorrhizal fungi

Available resources:

4 *Cortinarius* assemblies (ABYSS)
208 Agaricales single-copy orthologs
genes (SCOGs) & exonerate
Daicel Arbor Biosciences 

Resulting kit:

188 nuclear SCOGs
RPB1, RPB2, MCM7, GPD, & TEFI
~67.5 Kbp matrix
~20K 120-mer 2x tiled probes



Biological Journal of the Linnean Society, 2016, 117, 11–32. With 6 figures.

Tales from the crypt: genome mining from fungarium specimens improves resolution of the mushroom tree of life

BRYN T. M. DENTINGER^{1,2,*†}, ESTER GAYA^{1†}, HEATH O'BRIEN³, LAURA M. SUZ¹, ROBERT LACHLAN⁴, JORGE R. DÍAZ-VALDERRAMA⁵, RACHEL A. KOCH⁵ and M. CATHERINE AIME⁵

Fungal Diversity
<https://doi.org/10.1007/s13225-022-00499-9>



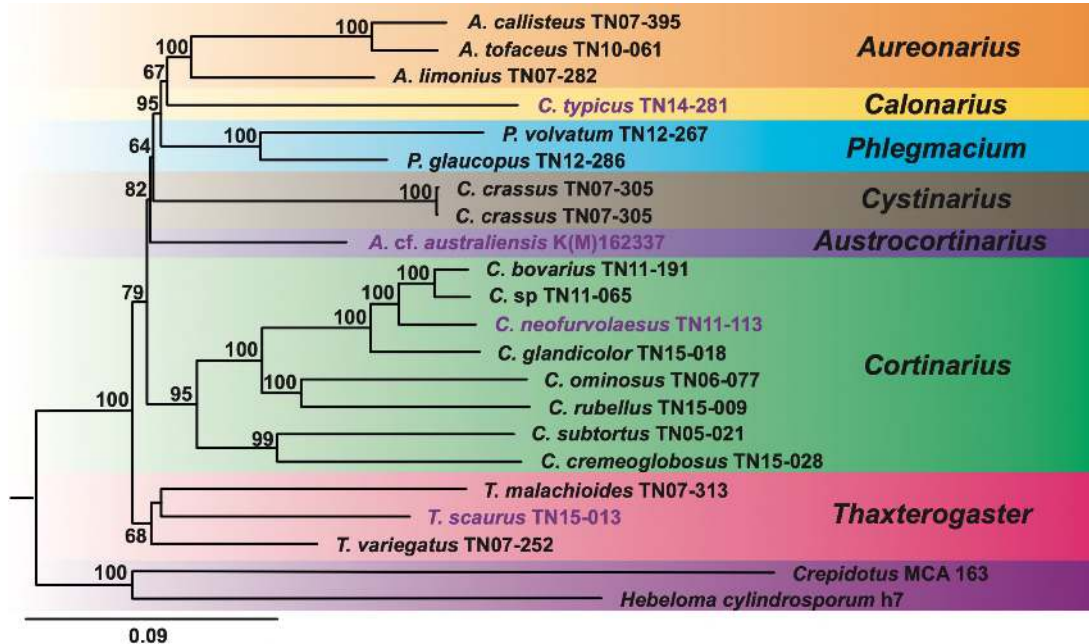
Taming the beast: a revised classification of *Cortinariaceae* based on genomic data

Kare Liimatainen¹  · Jan T. Kim²  · Lisa Pokorny^{1,3}  · Paul M. Kirk⁴  · Bryn Dentinger⁵  · Tuula Niskanen^{1,6} 

Received: 20 October 2021 / Accepted: 20 January 2022
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Cortinariaceae193





Angiosperms353

Classification: flowering plants, ~ 370K spp. (APG VI: 416 fams. & 64 orders)

Importance: agronomic, edible, timber, pulp, fiber, meds, ornamental, fuel, etc.

Available resources:

655 transcriptomes  **1KP**

20 genomes 

Hamming dist., k-medoids clustering

Daicel Arbor Biosciences 

Resulting kit:

353 nuclear SCOGs

~260,8 Kbp matrix

~75,2K 120-mer 3x tiled probes

Article

One thousand plant transcriptomes and the phylogenomics of green plants

<https://doi.org/10.1038/s41586-019-1693-2>

One Thousand Plant Transcriptomes Initiative

Syst. Biol. 68(4):594–606, 2019

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DOI:10.1093/sysbio/syy086
Advance Access publication December 10, 2018

A Universal Probe Set for Targeted Sequencing of 353 Nuclear Genes from Any Flowering Plant Designed Using k-Medoids Clustering

MATTHEW G. JOHNSON^{1,2,*}, LISA POKORNY³, STEVEN DODSWORTH^{3,4}, LAURA R. BOTIGUE^{3,5}, ROBYN S. COWAN³, ALISON DEVAULT⁶, WOLF L. EISERHARDT^{3,7}, NIROSHINI EPITAWALAGE³, FELIX FOREST³, JAN T. KIM³, JAMES H. LEEBENS-MACK⁸, ILIA J. LEITCH³, OLIVIER MAURIN³, DOUGLAS E. SOLTIS^{9,10}, PAMELA S. SOLTIS^{9,10}, GANE KA-SHU WONG^{11,12,13}, WILLIAM J. BAKER³, AND NORMAN J. WICKETT^{2,14}

¹Department of Biological Sciences, Texas Tech University, Lubbock, TX 79409, USA; ²Plant Science and Conservation, Chicago Botanic Garden, 1000 Lake Cook Road, Glenview, IL 60022, USA; ³Department of Comparative Plant and Fungal Biology, Royal Botanic Gardens, Kew, Richmond, Surrey TW9 3AE, UK; ⁴School of Life Sciences, University of Bedfordshire, University Square, Luton LU1 3JU, UK; ⁵Centre for Research in Agricultural Genomics, Campus UAB, Edifici CRAG, Bellaterra Cerdanyola del Vallès, 08193 Barcelona, Spain; ⁶Arbor Biosciences, 5840 Interface Dr, Suite 101, Ann Arbor, MI 48103, USA; ⁷Department of Bioscience, Aarhus University, 8000 Aarhus C, Denmark; ⁸Department of Plant Biology, University of Georgia, 2502 Miller Plant Sciences, Athens, GA 30602, USA; ⁹Department of Biology, University of Florida, 220 Bartman Hall, Gainesville, FL 32611-8525, USA; ¹⁰Florida Museum of Natural History, University of Florida, 3215 Hull Road, Gainesville, FL 32611-2710, USA; ¹¹BGI-Shenzhen, Beishan Industrial Zone, Yantian District, Shenzhen 518083, China; ¹²Department of Biological Sciences, University of Alberta, Edmonton, AB T6G 2E9, Canada; ¹³Department of Medicine, University of Alberta, Edmonton, AB T6G 2E1, Canada; and ¹⁴Program in Plant Biology and Conservation, Northwestern University, 2205 Tech Drive, Evanston, IL 60208, USA

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Matthew G. Johnson, Lisa Pokorny, Steven Dodsworth contributed equally to this article.



Angiosperms353



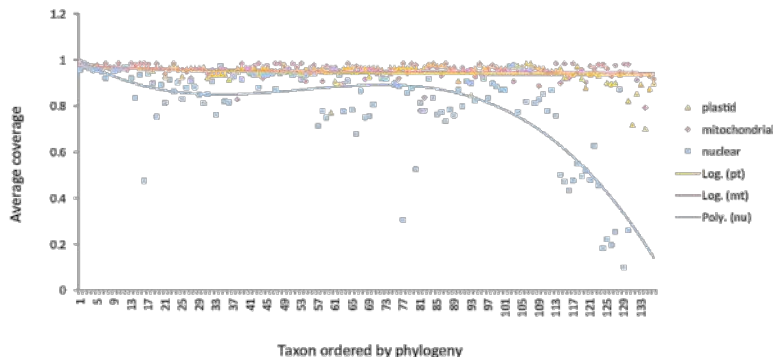
ARTICLE

<https://doi.org/10.1038/s41467-019-09454-w>

OPEN

Resolution of the ordinal phylogeny of mosses using targeted exons from organellar and nuclear genomes

Yang Liu^{1,2}, Matthew G. Johnson³, Cymon J. Cox⁴, Rafael Medina⁵, Nicolas Devos⁶, Alain Vanderpoorten⁷, Lars Hedenäs⁸, Neil E. Bell⁹, James R. Shevock¹⁰, Blanka Agüero⁶, Dietmar Quandt¹¹, Norman J. Wickett¹², A. Jonathan Shaw⁶ & Bernard Goffinet¹³



Article

One thousand plant transcriptomes and the phylogenomics of green plants

<https://doi.org/10.1038/s41586-019-1693-2>

One Thousand Plant Transcriptomes Initiative

Syst. Biol. 68(4):594–606, 2019

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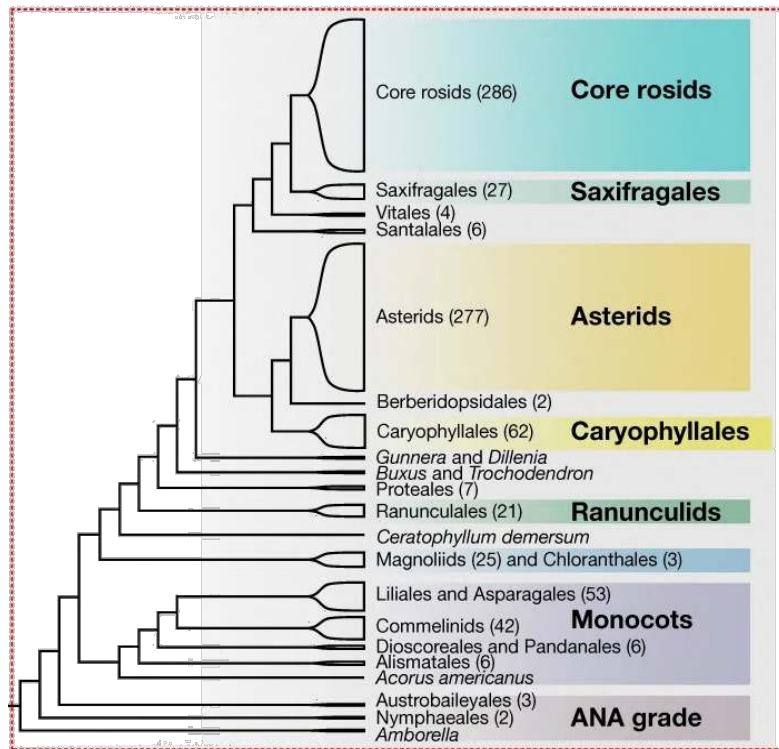
¹Department of Biological Sciences, Texas Tech University, Lubbock, TX 79409, USA; ²Plant Science and Conservation, Chicago Botanic Garden, 1000 Lake Cook Road, Glenview, IL 60022, USA; ³Department of Comparative Plant and Fungal Biology, Royal Botanic Gardens, Kew, Richmond, Surrey TW9 3AE, UK; ⁴School of Life Sciences, University of Bedfordshire, University Square, Luton LU1 3JU, UK; ⁵Centre for Research in Agricultural Genomics, Campus UAB, Edifici CRAG, Bellaterra Cerdanyola del Vallès, 08193 Barcelona, Spain; ⁶Arbor Biosciences, 5840 Interface Dr, Suite 101, Ann Arbor, MI 48103, USA; ⁷Department of Bioscience, Aarhus University, 8000 Aarhus C, Denmark; ⁸Department of Plant Biology, University of Georgia, 2502 Miller Plant Sciences, Athens, GA 30602, USA; ⁹Department of Biology, University of Florida, 220 Bartram Hall, Gainesville, FL 32611-8525, USA; ¹⁰Florida Museum of Natural History, University of Florida, 3215 Hull Road, Gainesville, FL 32611-2710, USA; ¹¹BGI-Shenzhen, Beishan Industrial Zone, Yantian District, Shenzhen 518083, China; ¹²Department of Biological Sciences, University of Alberta, Edmonton, AB T6G 2E9, Canada; ¹³Department of Medicine, University of Alberta, Edmonton, AB T6G 2E1, Canada; and ¹⁴Program in Plant Biology and Conservation, Northwestern University, 2205 Tech Drive, Evanston, IL 60208, USA

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Angiosperms353



Article

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<https://doi.org/10.1038/s41586-019-1693-2>

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Angiosperms353

ABCD1 target instance X and associated 120-mer probes



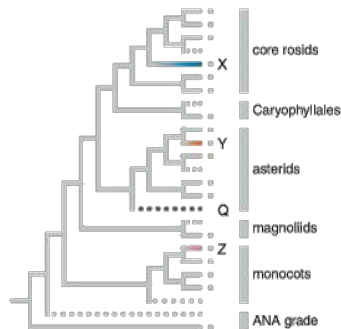
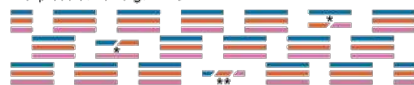
ABCD1 target instance Y and associated 120-mer probes



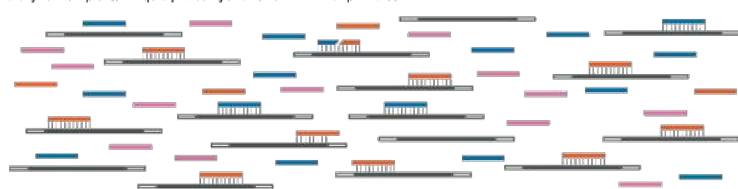
ABCD1 target instance Z and associated 120-mer probes



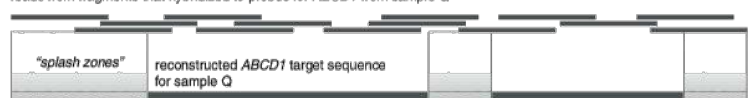
final probe set for target *ABCD1*



library for sample Q in liquid phase hybridization with final probe set



reads from fragments that hybridized to probes for *ABCD1* from sample Q



Syst. Biol. 68(4):594–606, 2019

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Matthew G. Johnson, Lisa Pokorny, Steven Dodsworth contributed equally to this article.



Angiosperms353

ABCD1 target instance X and associated 120-mer probes



ABCD1 target instance Y and associated 120-mer probes



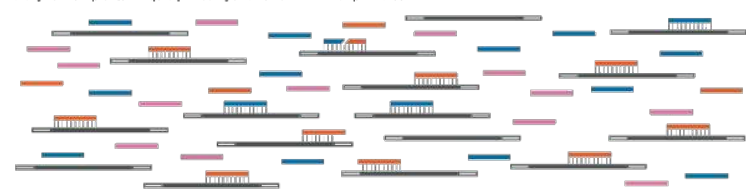
ABCD1 target instance Z and associated 120-mer probes



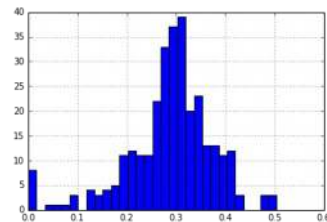
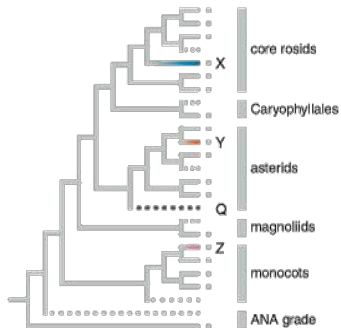
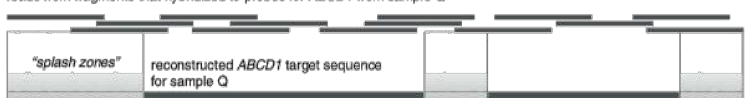
final probe set for target *ABCD1*



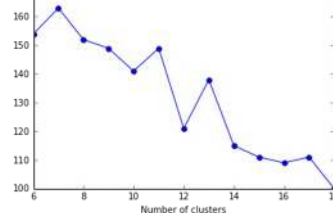
library for sample Q in liquid phase hybridization with final probe set



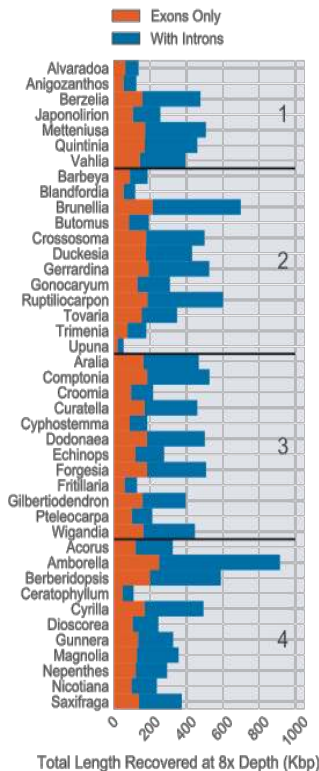
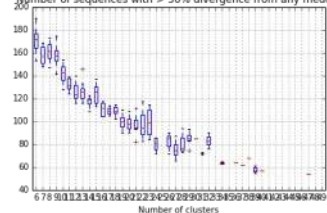
reads from fragments that hybridized to probes for *ABCD1* from sample Q



Number of sequences with > 30% divergence from any medoid



Boxplot grouped by *lim*Clusters





Angiosperms353



Genomics Synthetic Biology

myBaits Expert — Pre-designed Panels

myBaits Expert – Plant – Angiosperms-353

Home • Genomics • Targeted Sequencing • myBaits – Plant – Angiosperms-353 • myBaits Expert – Pre-designed Panels • myBaits Expert – Plant – Angiosperms-353



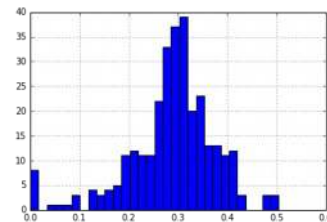
This new panel enriches 100's of single-copy genes orthologous across all angiosperms (flowering plants).

Combining the flexible hybridization power of in-solution target capture with an expertly selected set of orthologous locus sequences, this new probe set has been demonstrated to enrich hundreds of putatively single-copy protein-coding genes across a broad range of angiosperms (flowering plants). Probes were designed from 353 loci, each with 5-15 representative sequences from across all angiosperms which were selected using a novel "k-medoids clustering approach" to maximize taxonomic breadth of the design (Johnson, Pokorny, Dodsworth et al 2018, *Systematic Biology*).

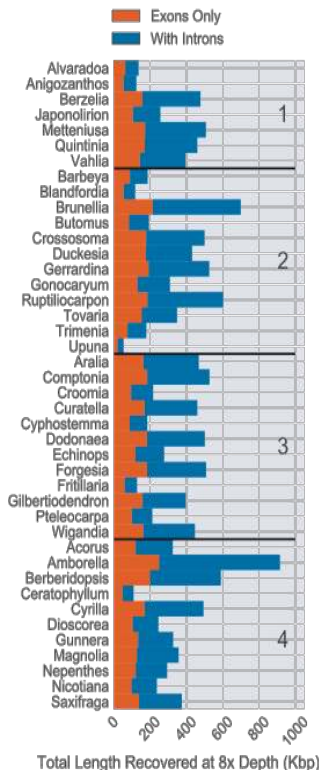
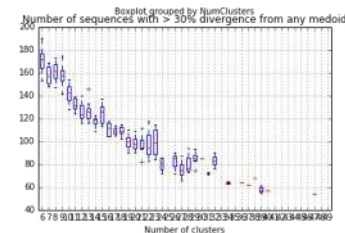
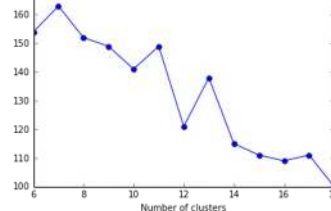
These probes are broadly applicable for phylogenetic research across all flowering plants, and the kit is available from Arbor Biosciences as an in-stock catalog kit available for immediate shipment at a low per-reaction cost. As with all myBaits kits, the Angiosperms353 panel is provided as a complete solution target capture kit, including buffers, blockers, and baits, along with an easy-to-use protocol. Or if you would prefer to outsource the work, our **myReads** NGS service expert scientists are available to perform library preparation, target capture, and sequencing for your entire project.

- Pre-designed Panel – Designed by expert plant geneticists
- Proven results – Demonstrated utility across broad taxonomic range
- Freely accessible pipelines – Simple, user-friendly data analysis
- Cost-effective – Use one bait set across multiple distantly-related taxa

Ordering Information	Size	Cat#	Qty	Price	
myBaits Expert Angiosperms 353 v1	8 Reactions	308108	1	\$1000	ADD TO CART
Available in-stock for immediate shipment.					



Number of sequences with > 30% divergence from any medoid





ProkarioKit

NIH National Library of Medicine
National Center for Biotechnology Information

Log in

Database of Clusters of Orthologous Genes (COGs)

Updated: March, 2022

COG DATABASE COG CATEGORIES PATHWAYS WEB SERVICES COG PROJECT

CONTACT

COGs stands for Clusters of Orthologous Genes. The database was initially created in 1997 (Tatusov et al., PMID: 9381173) followed by several updates, most recently in 2014 (Galperin et al., PMID: 25428365). The current update includes complete genomes of 1,187 bacteria and 122 archaea that map into 1,234 genera. The new features include ~250 updated COG annotations with corresponding references and PDB links, where available; new COGs for proteins involved in CRISPR-Cas immunity, sporulation, and photosynthesis, and the lists of COGs grouped by pathways and functional systems.

Search by:

COG Definition (COG0105 or just the number 105)
Any word in the COG name (polymerase)
Taxonomic Category (Mollicutes)
Organism name (Aciduliprofundum_boonei_T469)
Pathway (Arginine biosynthesis)
Assembly (GCA_D00091165.1)
Protein name: (prot:WP_011012300.1)
Gene Tag: (gene_tag:Haur_1857)

Search

Statistics

COGs	Genomic loci	Taxonomic Categories	Organisms	Protein IDs	COG symbols
4,877	3,456,089	37	1,309	3,213,255	3,828

Toward Automatic Reconstruction of a Highly Resolved Tree of Life

Francesca D. Ciccarelli,^{1,2,3,4} Tobias Doerks,^{1,4} Christian von Mering,¹ Christopher J. Creevey,¹ Berend Snij,² Peer Bork^{1,4,5}

We have developed an automatable procedure for reconstructing the tree of life with branch lengths comparable across all three domains. The tree has its basis in a concatenation of 31 orthologs occurring in 191 species with sequenced genomes. It revealed interdomain discrepancies in taxonomic classification. Systematic detection and subsequent exclusion of products of horizontal gene transfer increased phylogenetic resolution, allowing us to confirm accepted relationships and resolve disputed and preliminary classifications. For example, we place the phylum Acidobacteria as a sister group of β -Proteobacteria, support a Gram-positive origin of Bacteria, and suggest a thermophilic last universal common ancestor.

Reconstructing the phylogenetic relationships among all living organisms is one of the fundamental challenges in biology. Numerous attempts to derive a tree of life using various methods have been published [for a review, see (1)], and its principal existence has been questioned recently (2, 3). Moreover, even under the assumption of a tree of life, numerous groupings and taxonomic entities still remain heavily debated, and the advent of molecular and genomic data has increased the variety of classifications rather than reducing the problems (4).

Theoretical and practical limits to reconstructing a tree of life have been put forward, such as the insufficient amount of discriminating characters available, even in information-rich genomic data sets (4), and the computing resources required to cope with large numbers of species (1). Furthermore, there are factors that hamper accurate reconstruction of phylogenetic trees regardless of the methods used, such as sampling biases of species included (5) and dilution of phylogenetic signal by horizontal gene transfer (HGT) (6), the

¹European Molecular Biology Laboratory, Meyerhofstrasse 1, 60402 Heidelberg, Germany; ²Department of Evolutionary Genetics, European Institute of Oncology, Via E. Fermi 485, 20145 Milan, Italy; ³Institute of Molecular Biology, University of Zurich, Winterthurerstrasse 190, 8057 Zurich, Switzerland; ⁴Center for Molecular and Biomolecular Informatics, Max Planck Institute for Molecular Evolutionary Biology, Am Fassberg 11, 42205 Essen, Germany; ⁵Department of Biology, University of California, San Diego, La Jolla, CA 92037, USA

These authors contributed equally to this work. To whom correspondence should be addressed: E-mail: prokariokit@ebi.ac.uk

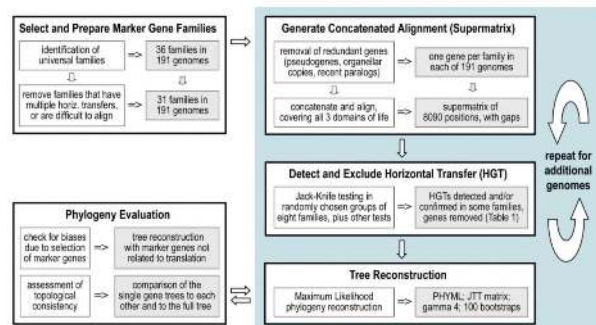
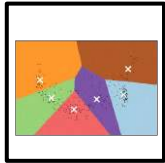


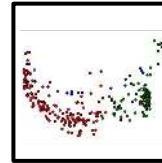
Fig. 1. Overview of the procedure. The white boxes represent the major steps for building the pan-domain phylogeny presented here. Steps in gray represent automatable parts of the procedure that need to be carried out for including further species. For the 31 clusters of orthologous groups (COGs) used in the analysis, we manually derived 31 orthologs by removing mitochondrial and chloroplast paralogues from corresponding multiple alignments. We built domain-specific alignments by using corresponding protein encoded by the 31 orthologs and aligned the resulting profiles. With this procedure, we maximized the number of positions of the global alignment and reduced the number of misaligned residues. For a detailed description of the method, see (8).

Clustering Algorithms



Centroid

Proximity, e.g., K-means, K-medoids



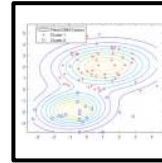
Graph

Distance, e.g., spectral (affinity propagation)



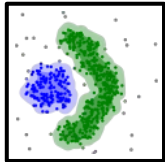
Connectivity

Agglomerative & Divisive (Hierarchical)



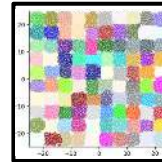
Distribution

Belonging, Gaussian Mixture Models



Density

DBSCAN, OPTICS, etc.



Compression

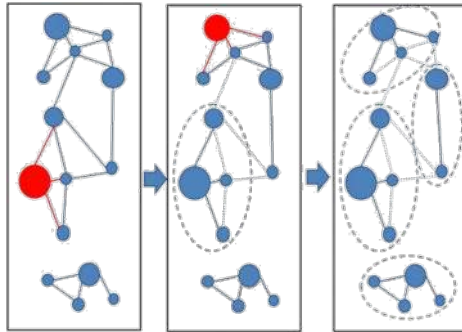
Dim. Transformation, e.g., BIRCH

Biological Sequence Clustering



CD-HIT

Greedy Incremental Clustering

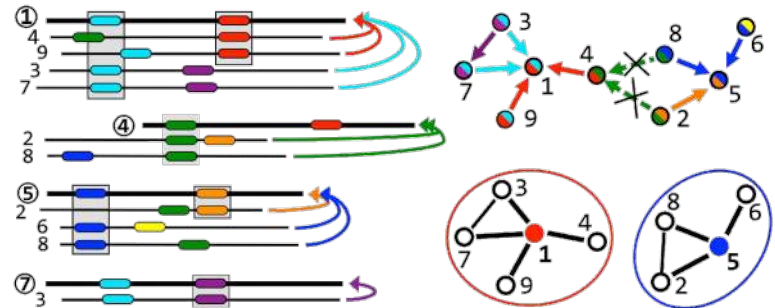


<https://github.com/soedinglab/mmseqs2/wiki#clustering-modes>



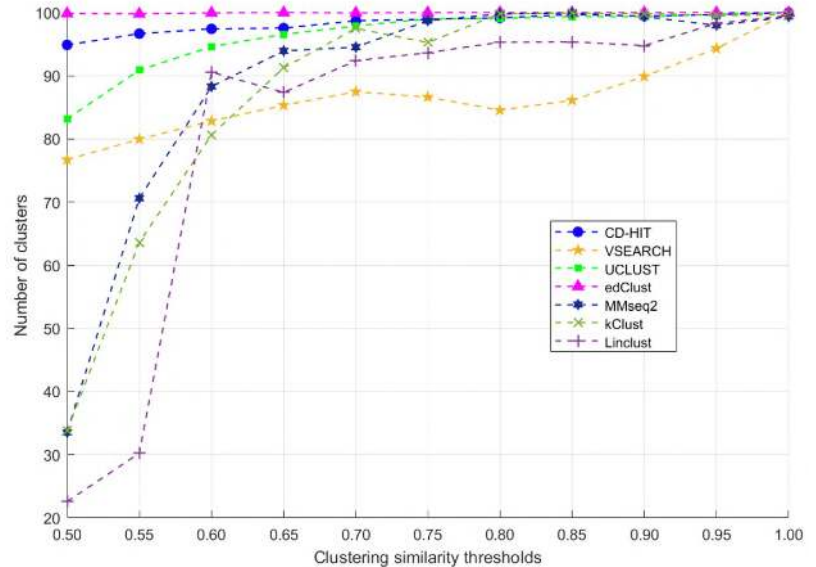
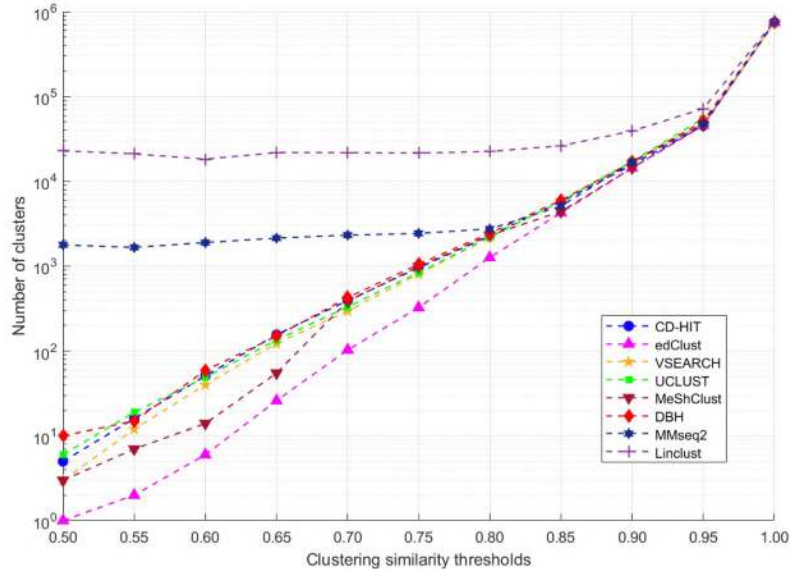
MMseqs2

Linear Time Clustering



<https://github.com/soedinglab/mmseqs2/wiki#linear-time-clustering-using-mmseqs-linclust>

Biological Sequence Clustering



Figs. 3 & 7. Wei Z.-G. et al. 2023. Comparison of Methods for Biological Sequence Clustering. *IEEE/ACM Trans. Comput. Biol. Bioinform.* 20(5): 2874–2888.

Tree-Based Clustering

PLOS ONE

RESEARCH ARTICLE

TreeCluster: Clustering biological sequences using phylogenetic trees

Metin Balaban¹, Niema Moshiri¹, Uyen Mai², Xingfan Jia³, Siavash Mirarab^{4*}

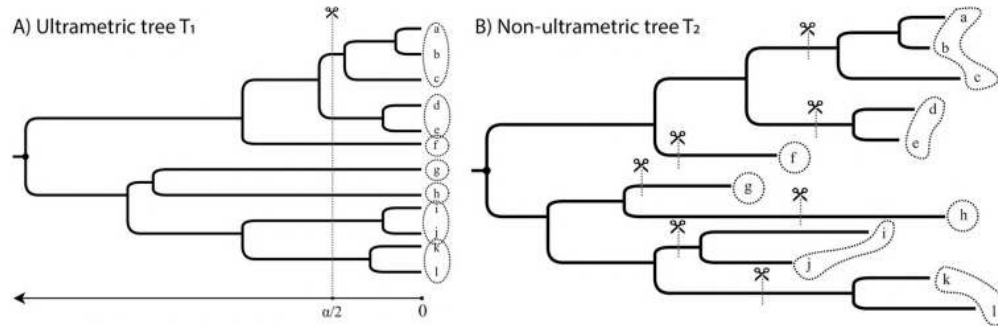


Fig 1. When the phylogenetic tree is ultrametric, clustering is trivial. For a threshold α , cut the tree at $\frac{\alpha}{2}$ height (A). When the tree is not ultrametric, it is not obvious how to cluster leaves (B). In both cases, a set of cut edges defines a clustering.

<https://doi.org/10.1371/journal.pone.0221068.g001>

Table of contents

01

Genome Subsampling

An Introduction to TCS
Approaches

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Probe Set Design

Custom & Universal Kits

03

Implementing TCS

Wet Lab Workflow &
Dry Lab Pipeline

04

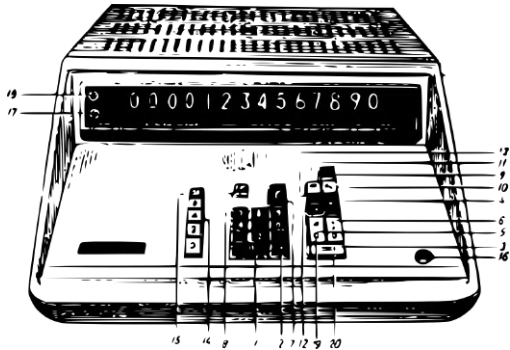
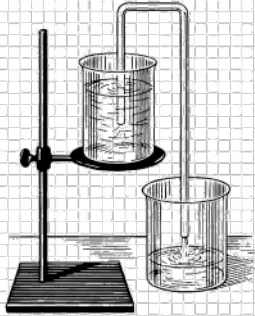
Biodiversity Genomics

Natural History Collections

Implementing TCS

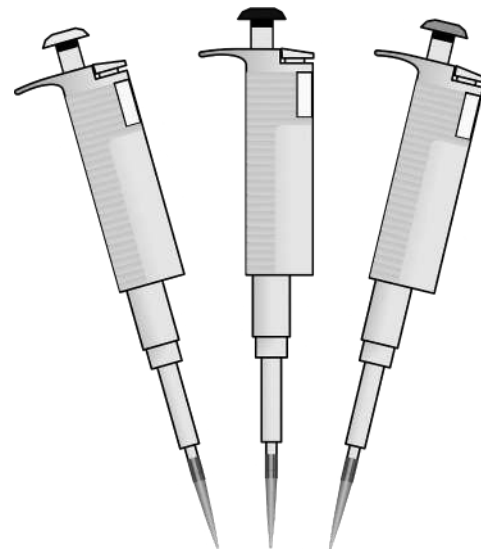
Wet Lab Workflow & Dry Lab Pipeline

03



Wet Lab Workflow

Generating TCS data



Wet Lab Workflow

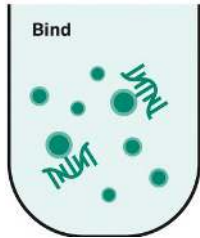
DNA Extraction (modified CTAB) & Purification
(cleaning w/ magnetic beads)

Tissue
Template

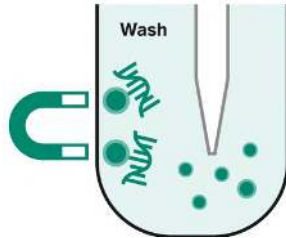


Purified DNA
Template

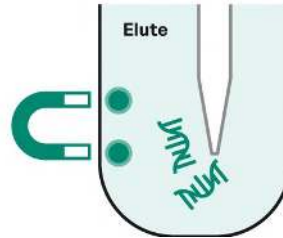
Magnetic particles
are added to sample
and bind to target
molecule



Magnetic particles
are captured and
remainder of sample
is washed away



Target molecule is
released from
magnetic particles
for further analysis



Dioscorea tomentosa, 1796

Wet Lab Workflow

DNA Extraction (modified CTAB) & Purification
(cleaning w/ magnetic beads)

Genomic Library Preparation (commercially
available kits for dual-indexed sequencing)

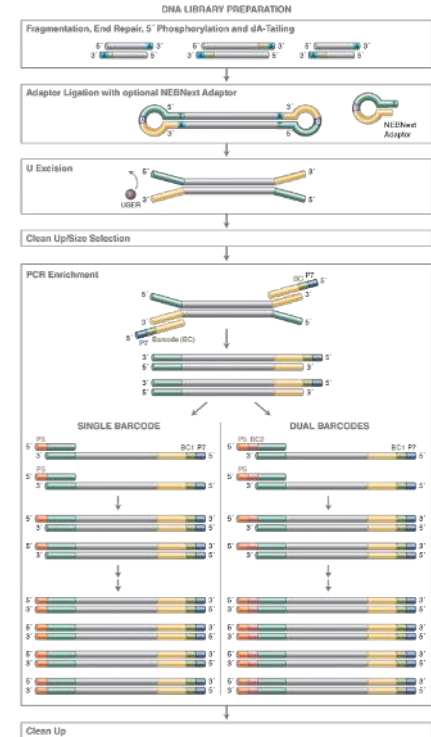


Tissue
Template



Purified DNA
Template

Dual-indexed
Genomic Libraries



Wet Lab Workflow

DNA Extraction (modified CTAB) & Purification
(cleaning w/ magnetic beads)

Genomic Library Preparation (commercially
available kits for dual-indexed sequencing)

Hybridization (biotinylated probes), Capture
(streptavidin-coated beads) & Enrichment (PCR)



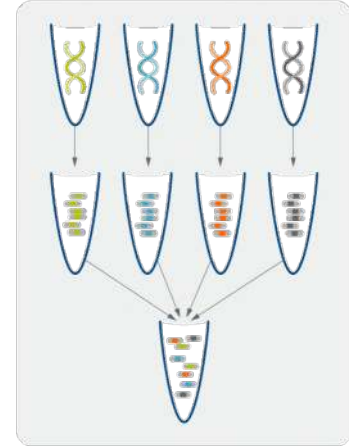
Tissue
Template



Purified DNA
Template

Dual-indexed
Genomic Libraries

Target-enriched
Genomic Libraries



Wet Lab Workflow

DNA Extraction (modified CTAB) & Purification
(cleaning w/ magnetic beads)

Genomic Library Preparation (commercially
available kits for dual-indexed sequencing)

Hybridization (biotinylated probes), Capture
(streptavidin-coated beads) & Enrichment (PCR)

High-Throughput Sequencing (HTS, multiplexed
enriched dual-indexed libraries)

Tissue
Template



Purified DNA
Template

Dual-indexed
Genomic Libraries

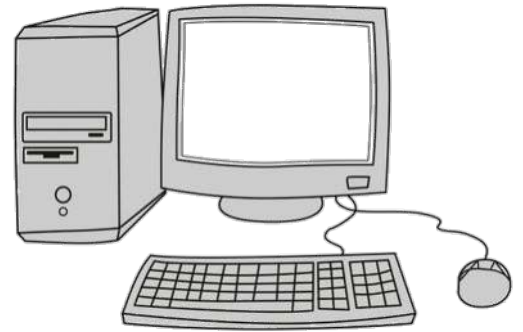
Target-enriched
Genomic Libraries

Raw
Reads



Dry Lab Pipeline

Processing TCS data



Dry Lab Pipeline

Raw
Reads

Cut



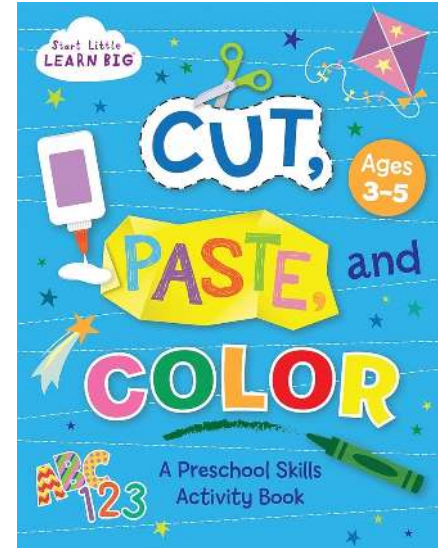
Paste



Color



Target
sequences



Dry Lab Pipeline

Raw
Reads

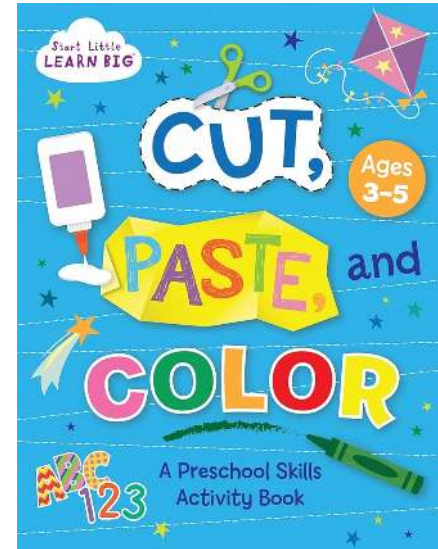
FastQC | MultiQC & Trimmomatic | fastp

BLAST | BWA | DIAMOND

SPAdes | exon(intron)erate


HybPiper

Target
sequences

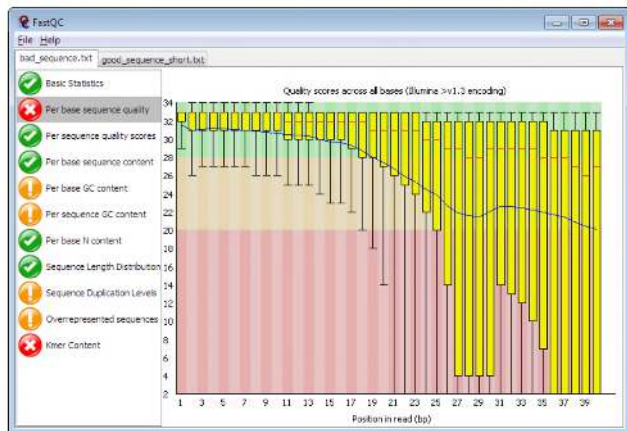


FastQC



FastQC

FastQC is a program designed to spot potential problems in high throughput sequencing datasets. It runs a set of analyses on one or more raw sequence files in fastq or bam format and produces a report which summarises the results.



MultiQC



Aggregate bioinformatics results across many samples into a single report

Find [documentation](http://multiqc.info) and [example reports](http://multiqc.info) at <http://multiqc.info>

MultiQC v1.10 bioconductor v1.10 DOI: 10.1093/bioinformatics/btq154

MultiQC is a tool to create a single report with interactive plots for multiple bioinformatics analyses across many samples.

Reports are generated by scanning given directories for recognised log files. These are parsed and a single HTML report is generated summarising the statistics for all logs found. MultiQC reports can describe multiple analysis steps and large numbers of samples within a single plot, and multiple analysis tools making it ideal for routine fast quality control.

A very large number of Bioinformatics tools are supported by MultiQC. Please see the MultiQC website for a [complete list](http://multiqc.info). MultiQC can also easily parse data from custom scripts, if correctly formatted / configured - a feature called [Custom Content](http://multiqc.info).

trimmomatic

BIOINFORMATICS ORIGINAL PAPER

Vol. 30 no. 15 2014, pages 2114–2120
doi:10.1093/bioinformatics/btu170

Genome analysis

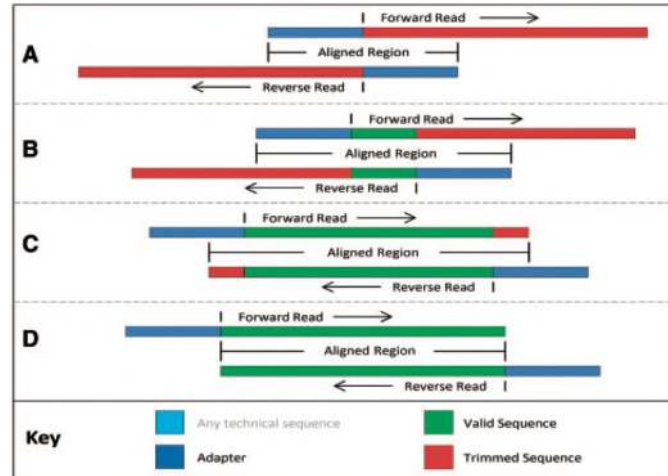
Advance Access publication April 1, 2014

Trimmomatic: a flexible trimmer for Illumina sequence data

Anthony M. Bolger^{1,2}, Marc Lohse¹ and Bjoern Usadel^{2,3,*}

¹Department Metabolic Networks, Max Planck Institute of Molecular Plant Physiology, Am Mühlenberg 1, 14476 Golm, ²Institut für Biologie I, RWTH Aachen, Worringer Weg 3, 52074 Aachen and ³Institute of Bio- and Geosciences: Plant Sciences, Forschungszentrum Jülich, Leo-Brandt-Straße, 52425 Jülich, Germany

Associate Editor: Inanc Biral

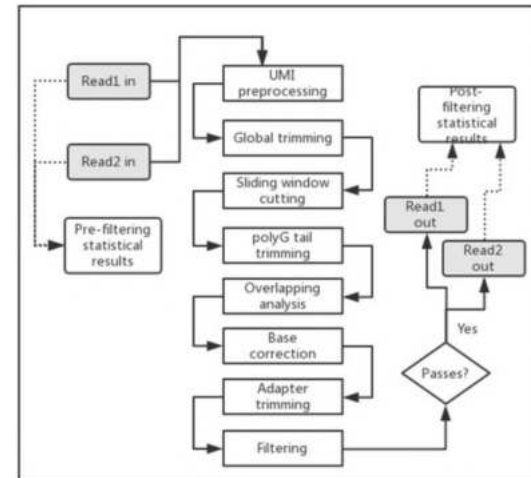


fastp

Bioinformatics, 34, 2018, i884–i890
doi: 10.1093/bioinformatics/bty560
ECCB 2018

fastp: an ultra-fast all-in-one FASTQ preprocessor

Shifu Chen^{1,2,*}, Yanqing Zhou¹, Yaru Chen¹ and Jia Gu²



HybPiper



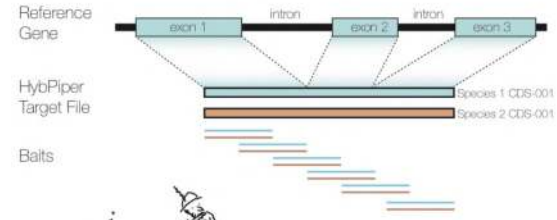
Applications in Plant Sciences 2016 4(7): 1600016

SOFTWARE NOTE

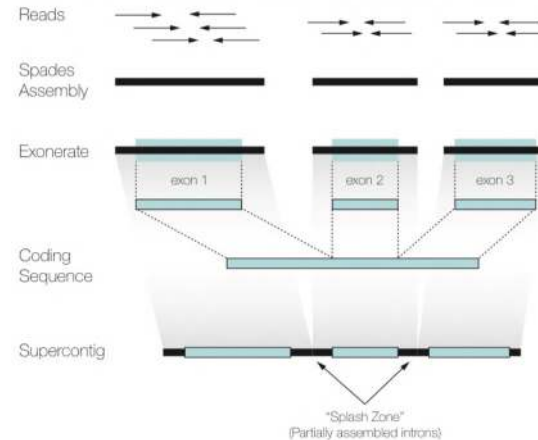
HYBPIPER: EXTRACTING CODING SEQUENCE AND INTRONS FOR PHYLOGENETICS FROM HIGH-THROUGHPUT SEQUENCING READS USING TARGET ENRICHMENT¹

MATTHEW G. JOHNSON^{2,6}, ELLIOT M. GARDNER^{2,3}, YANG LIU⁴, RAFAEL MEDINA⁴,
BERNARD GOFFINET⁴, A. JONATHAN SHAW⁵, NYREE J. C. ZERBICA^{2,3}, AND NORMAN J. WICKETT^{2,3}

Target file and bait design (pre-HybPiper)



HybPiper



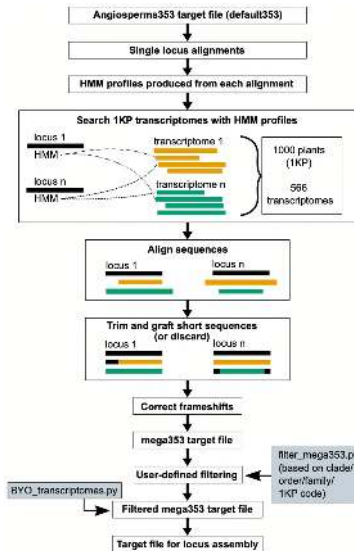
HybPiper

INVITED SPECIAL ARTICLE

For the Special Issue: Exploring Angiosperms353: A Universal Toolkit for Flowering Plant Phylogenomics

New targets acquired: Improving locus recovery from the Angiosperms353 probe set

Todd G. B. McLay^{1,2,3}, Joanne L. Birch¹, Bee F. Gunn¹, Weixuan Ning¹, Jennifer A. Tate¹, Lars Nauheimer¹, Elizabeth M. Joyce¹, Laitha Simpson^{1,2}, Alexander N. Schmidt-Lebuhn^{1,2}, William J. Baker¹, Felix Forest¹, and Chris J. Jackson¹



Received: 21 September 2022

Accepted: 30 January 2023

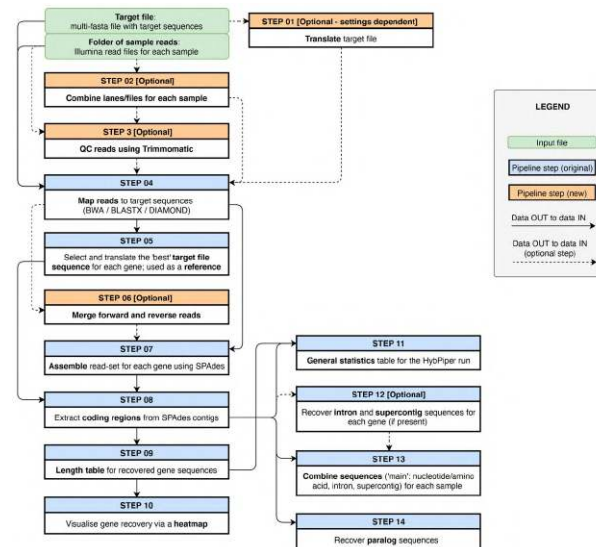
DOI: 10.1002/aps3.11532

SOFTWARE NOTE



hybpiper-nf and paragone-nf: Containerization and additional options for target capture assembly and paralog resolution

Chris Jackson¹ | Todd McLay^{1,2,3} | Alexander N. Schmidt-Lebuhn²



PHYLUC

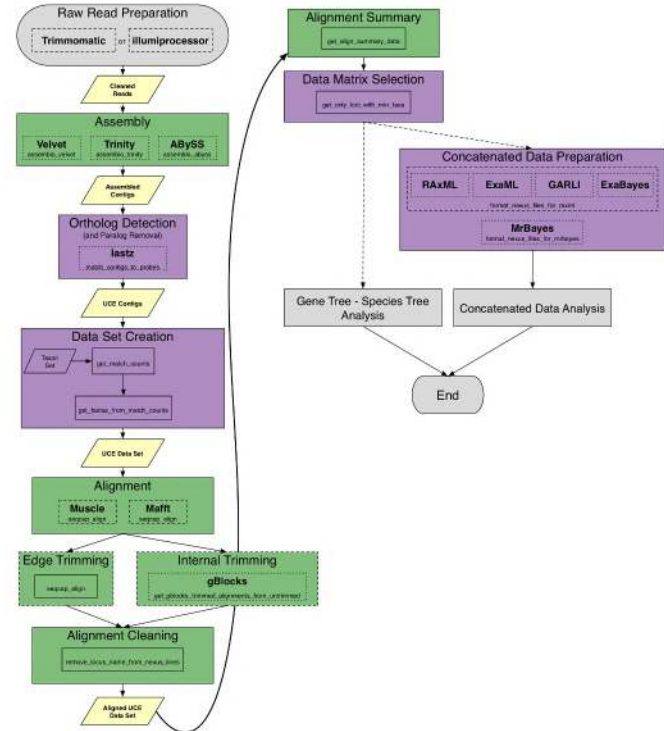
Bioinformatics, 32(5), 2016, 786–788
doi: 10.1093/bioinformatics/btv646
Advance Access Publication Date: 2 November 2015
Applications Note

OXFORD

Phylogenetics

PHYLUC is a software package for the analysis of conserved genomic loci

Brant C. Faircloth

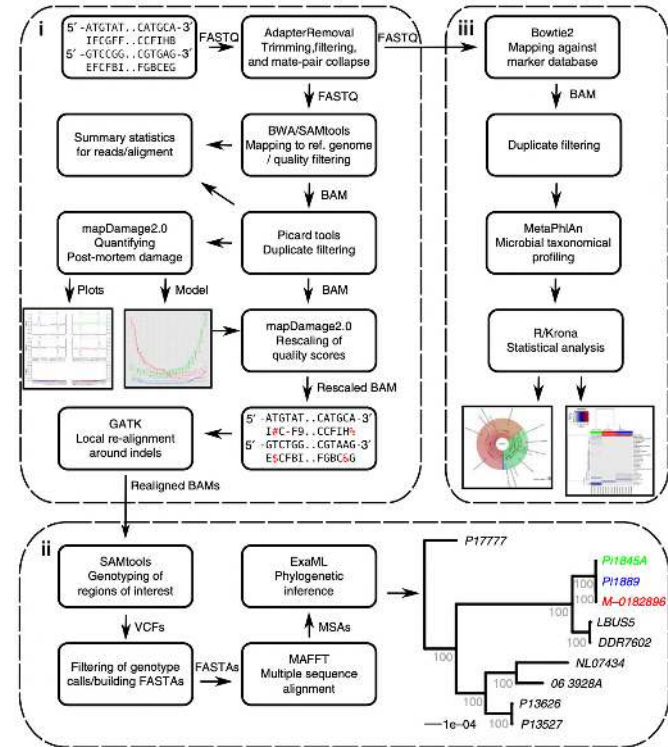


PALEOMIX

PROTOCOL

Characterization of ancient and modern genomes by SNP detection and phylogenomic and metagenomic analysis using PALEOMIX

Mikkel Schubert¹, Luca Ermini¹, Clio Der Sarkissian¹, Hákon Jónsson¹, Aurélien Ginolhac¹, Robert Schaefer², Michael D Martin¹, Ruth Fernández¹, Martin Kircher³, Molly McCue⁴, Eske Willerslev¹ & Ludovic Orlando¹

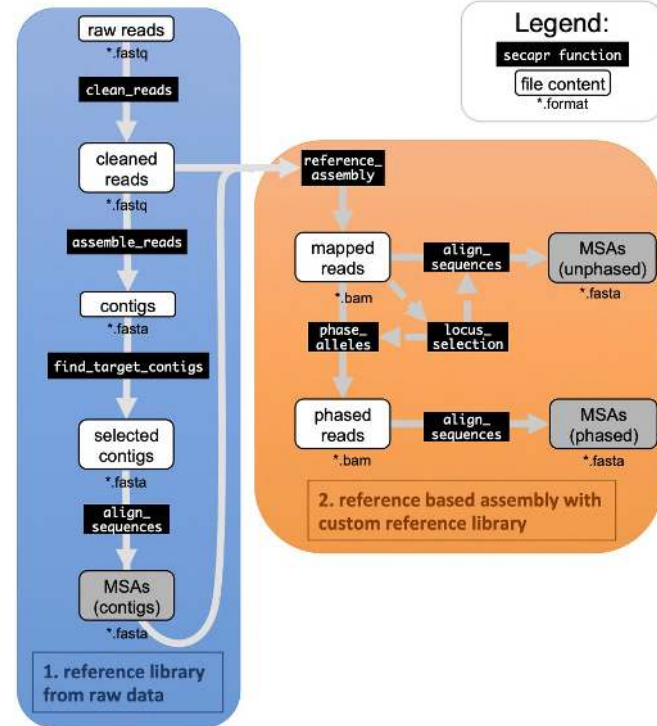




SECAPR

SECAPR—a bioinformatics pipeline for the rapid and user-friendly processing of targeted enriched Illumina sequences, from raw reads to alignments

Tobias Andermann^{1,2}, Ángela Cano^{2,3}, Alexander Zizka^{1,2},
Christine Bacon^{1,2} and Alexandre Antonelli^{1,2,4,5}



HybPhyloMaker

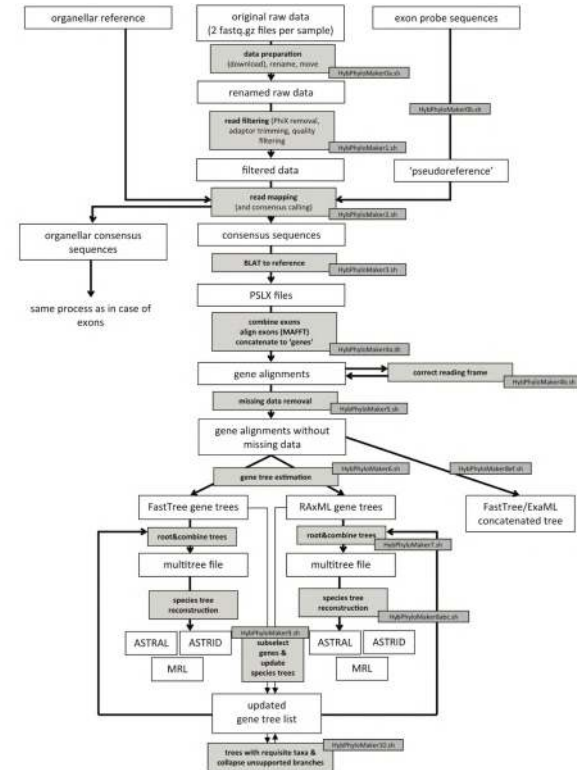
Evolutionary Bioinformatics
Volume 14, January–December 2018
© The Author(s) 2018, Article Reuse Guidelines
<https://doi.org/10.1177/1176934317742613>



Short Report

HybPhyloMaker: Target Enrichment Data Analysis From Raw Reads to Species Trees

Tomáš Fér¹ and Roswitha E Schmickl²



PAFTools

Syst. Biol. 71(2):301–319, 2022

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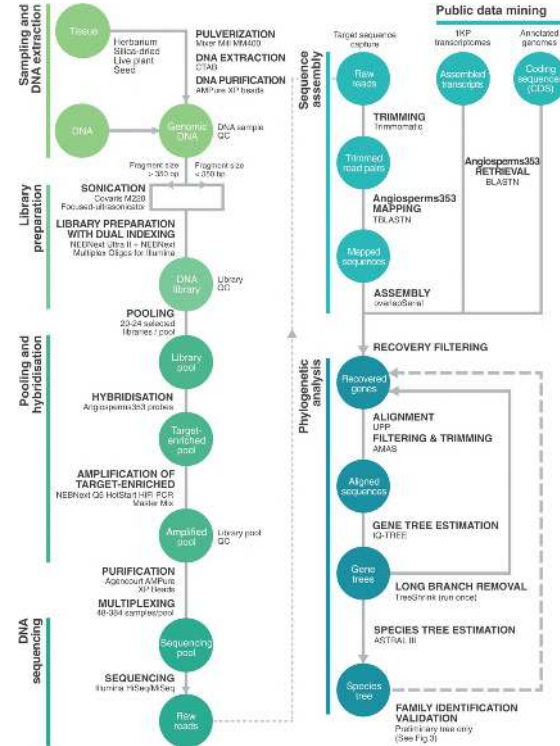
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DOI:10.1093/sysbio/syab055

Advance Access publication May 13, 2021

A Comprehensive Phylogenomic Platform for Exploring the Angiosperm Tree of Life

WILLIAM J. BAKER^{1,*}, PAUL BAILEY¹, VANESSA BARKER¹, ABIGAIL BARKER¹, SIDONIE BELLOT¹, DAVID BISHOP¹, LAURA R. BOTICOURT^{1,2}, GRACE BREWER¹, TOM CARRUTHERS¹, JAMES J. CLARKSON¹, JEFFREY COOK¹, ROBYN S. COWAN¹, STEVEN DOUSWORTH^{1,3}, NIROSHINI EPIPAWALAGE¹, ELAINE FRANCOSO¹, BERKA GALLEGO¹, MATTHEW G. JOHNSON⁴, JAN T. KIM^{1,5}, KEVIN LEEMPOEL¹, OLIVIER MAURIN¹, CATHERINE MCGINNIE¹, LISA POKORNY^{1,6}, SHYAMALI ROY¹, MALCOLM STONE¹, EDUARDO TOLEDO¹, NORMAN J. WICKETT⁷, ALEXANDRE R. ZUNTINI¹, WOLF L. EISENHARDT^{1,8}, PAUL J. KERSEY¹, ILIJIA J. LEITCH¹, AND FELIX FOREST¹



CAPTUS

bioRxiv preprint doi: <https://doi.org/10.1101/2023.10.27.564367>; this version posted November 1, 2023. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

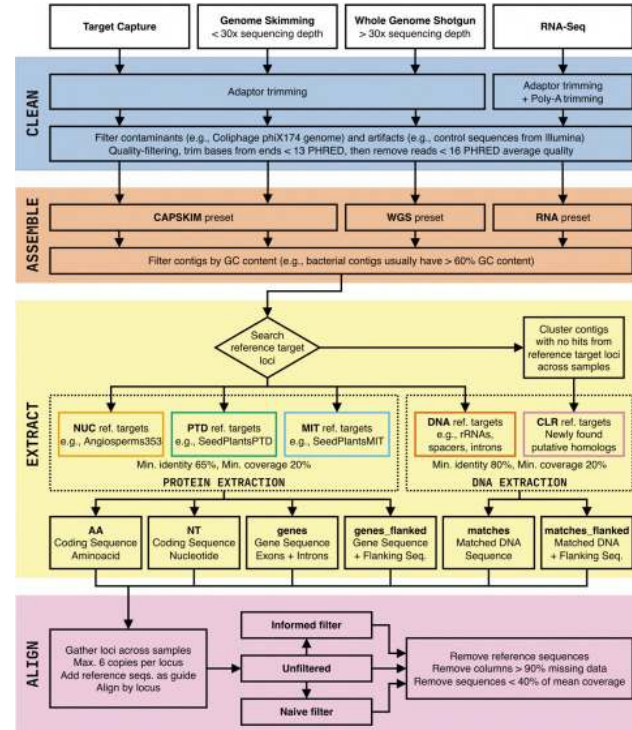
A novel phylogenomics pipeline reveals complex pattern of reticulate evolution in

Cucurbitales

Running Title: CAPTUS: a novel pipeline for phylogenomics

Edgardo M. Ortiz^{1*}, Alina Höwener^{1,2}, Gentaro Shigita¹, Mustafa Raza¹, Olivier Maurin³,

Alexandre Zuntini³, Félix Forest³, William J. Baker³ & Hanno Schaefer¹



Dry Lab Pipeline

Raw
Reads

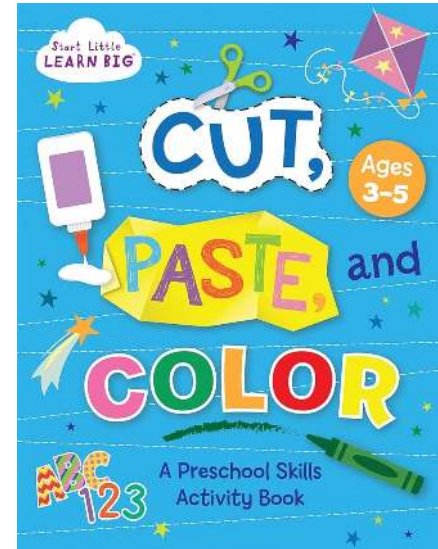
FastQC | MultiQC & Trimmomatic | fastp

BLAST | BWA | DIAMOND

SPAdes | exon(intron)erate


HybPiper

Target
sequences



Dry Lab Pipeline

Target
Sequences

Wash

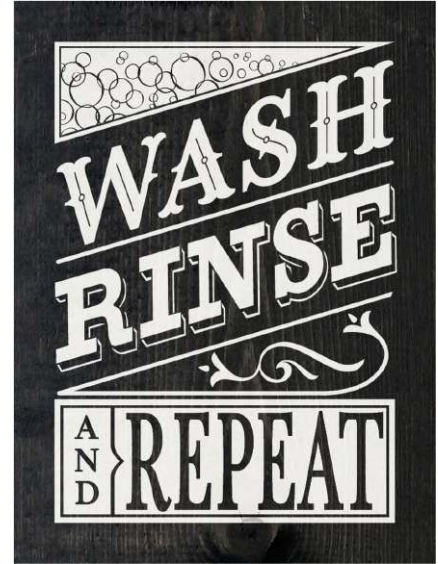


Rinse

Repeat



Aligned
Matrices



Dry Lab Pipeline

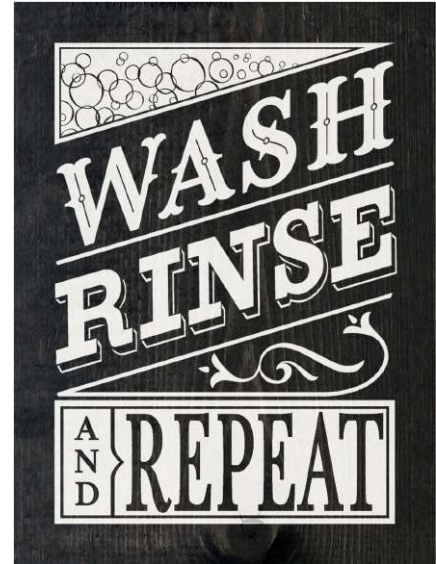
Target
Sequences

MAFFT | MUSCLE | PASTA | UPP

AMAS | PhyKIT | trimAl | ClipKIT

TreeShrink | PhylteR

Aligned
Matrices



MAFFT

© 2002 Oxford University Press

Nucleic Acids Research, 2002, Vol. 30 No. 14 3059–3066

MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform

Kazutaka Katoh, Kazuharu Misawa¹, Kei-ichi Kuma and Takashi Miyata*

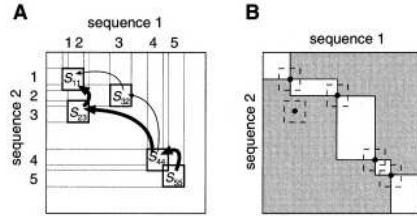
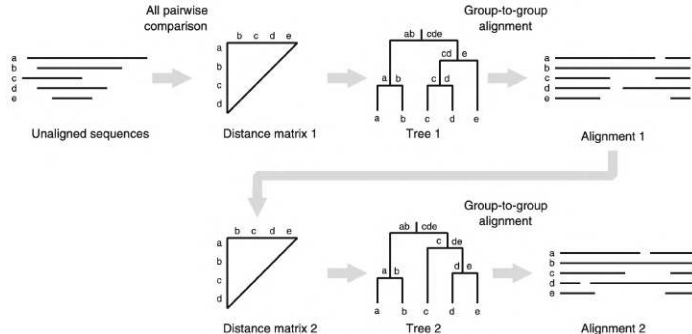


Figure 2. (A) An example of the segment-level DP; (B) Reducing the area for DP on a homology matrix.



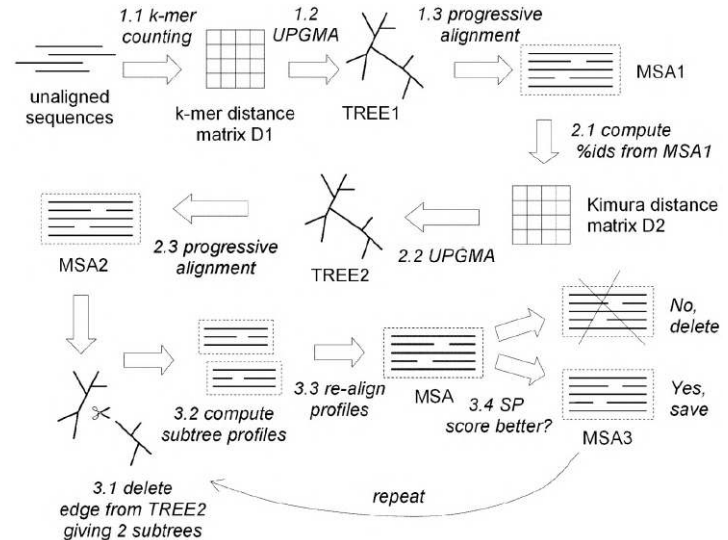
MUSCLE

1792–1797 Nucleic Acids Research, 2004, Vol. 32, No. 5

DOI: 10.1093/nar/gkh340

MUSCLE: multiple sequence alignment with high accuracy and high throughput

Robert C. Edgar*

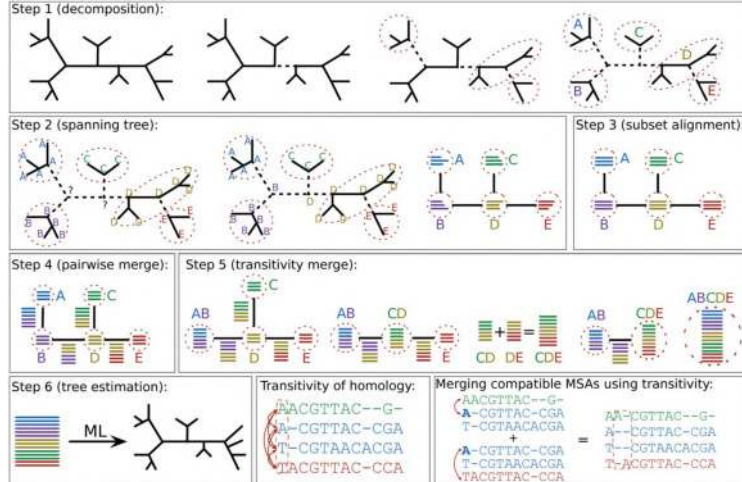


PASTA

JOURNAL OF COMPUTATIONAL BIOLOGY
Volume 22, Number 5, 2015
© Mary Ann Liebert, Inc.
Pp. 377–386
DOI: 10.1089/cmb.2014.0156

PASTA: Ultra-Large Multiple Sequence Alignment for Nucleotide and Amino-Acid Sequences

SIYAVASH MIRARAB¹, NAM NGUYEN¹, SHENG GUO², LI-SAN WANG,³
JUNHYONG KIM,³ and TANDY WARNOW^{1,4}



UPP

Nguyen et al. *Genome Biology* (2015) 16:124
DOI 10.1186/s13059-015-0688-z



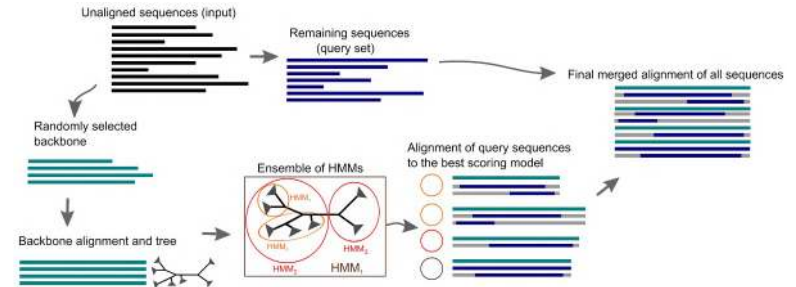
METHOD

Open Access



Ultra-large alignments using phylogeny-aware profiles

Nam-phuong D. Nguyen¹, Siavash Mirarab², Keerthana Kumar² and Tandy Warnow^{1,3,4*}



AMAS

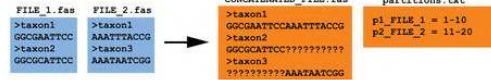
PeerJ

AMAS: a fast tool for alignment manipulation and computing of summary statistics

Marek L. Borowiec

A: concatenation

AMAS concat -i FILE_1.fas FILE_2.fas -f fasta -d dna --concat-out CONCATENATED_FILE.fas



B: format conversion

AMAS convert -i FILE_1.fas FILE_2.fas -f fasta -d dna --out-format phylip



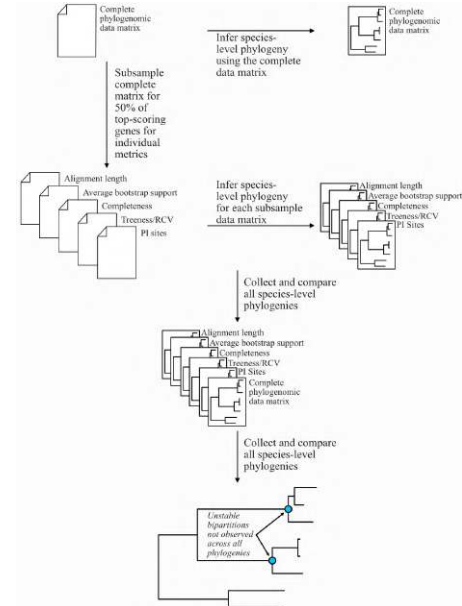
Function	AMAS	FASconCAT-G	Phyutility
Input formats	fasta phylip nexus	clustal fasta phylip nexus	fasta nexus
Concatenation	yes	yes	yes
Splitting or site extraction	yes	yes	yes (gaps only)
Summary statistics	yes	yes	no
Replicate alignments	yes	no	no
Taxon removal	yes	no	no
Translation	no	yes	no
RY coding	no	yes	no
Consensus sequences	no	yes	no
NCBI interactions	no	no	yes
Tree manipulations	no	no	yes

PhyKIT

Phylogenetics

PhyKIT: a broadly applicable UNIX shell toolkit for processing and analyzing phylogenomic data

Jacob L. Steenwyk^{1,*}, Thomas J. Buida III², Abigail L. Labella², Yuaning Li², Xing-Xing Shen³, & Antonis Rokas^{1,4,*}



phyutility

BIOINFORMATICS APPLICATIONS NOTE

Vol. 24 no. 5 2008, pages 715–716
doi:10.1093/bioinformatics/btm619

Phylogenetics

Phyutility: a phyloinformatics tool for trees, alignments and molecular data

Stephen A. Smith^{1,*} and Casey W. Dunn²

3 Tree Functions

- 3.1 Rerooting
 - 3.1.1 Examples
- 3.2 Pruning
 - 3.2.1 Examples
- 3.3 Type conversion
 - 3.3.1 Examples
- 3.4 Consensus
 - 3.4.1 Examples
- 3.5 Leaf stability
 - 3.5.1 Examples
- 3.6 Branch Attachment Frequency
 - 3.6.1 Examples
- 3.7 Tree support
 - 3.7.1 Examples
- 3.8 Thinning trees
 - 3.8.1 Examples

4 Data matrix functions

- 4.1 Concatenate
 - 4.1.1 Examples
- 4.2 GenBank Parsing
 - 4.2.1 Examples
- 4.3 Trimming sites
 - 4.3.1 Examples
- 4.4 Searching NCBI
 - 4.4.1 Examples
- 4.5 Fetching Sequences from NCBI
 - 4.5.1 Examples

FASconCAT-G

Kück and Longo *Frontiers in Zoology* 2014, **11**:81
<http://www.frontiersinzoology.com/content/11/1/81>



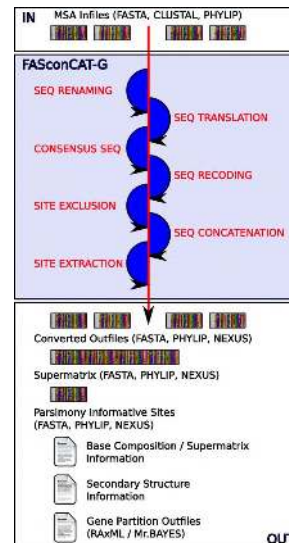
FRONTIERS IN ZOOLOGY

METHODOLOGY

Open Access

FASconCAT-G: extensive functions for multiple sequence alignment preparations concerning phylogenetic studies

Patrick Kück^{1*} and Gary C. Longo²



trimAl

BIOINFORMATICS APPLICATIONS NOTE

Vol. 25 no. 15 2009, pages 1972–1973
doi:10.1093/bioinformatics/btp348

Phylogenetics

trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses

Salvador Capella-Gutiérrez, José M. Silla-Martínez and Toni Gabaldón*

```
trimAl -in example1 -out output1 -htmlout output1.html -gt 1
```

Selected Residue / Sequence

Deleted Residue / Sequence

	10	20	30	40	50	60
Sp8	-----GLGKV	---IVY	-GIVLGTKS	-DQFSNVVWL	----	FPWNGQLQHMMGII
Sp17	-----FAYTAPD	---LLLIGFLLKTVA	-T-FG	-DTWF	----	QLWQGLDLNKMVPF
Sp10	-----DPAVL	---FV	-IMLGTIT	-K-FS	-SEWF	FAWLGLEINMMVII
Sp26	AAAAA	---ALL	---TYL	-GLFLGTDY	----	EN--FAAAAAANAWLGLEINMAQI
Sp33	-----PTIL	---	NIA	-GLHMETDI	-N-FS	-LAWF--QAWGGLEINKQAIL
Sp6	-----ASGAI	---	LTL	-GIYLFITLC	-AVIS	-VSWY----LAWLGLEINMMII

```
trimAl -in example1 -out output3 -htmlout output3.html -gt 0.8 -st 0.001 -cons 60
```

Selected Residue / Sequence

Deleted Residue / Sequence

	10	20	30	40	50	60
Sp8	-----GLGKV	---IVY	-GIVLGTKS	-DQFSNVVWL	----	FPWNGQLQHMMGII
Sp17	-----FAYTAPD	---LLLIGFLLKTVA	-T-FG	-DTWF	----	QLWQGLDLNKMVPF
Sp10	-----DPAVL	---FV	-IMLGTIT	-K-FS	-SEWF	FAWLGLEINMMVII
Sp26	AAAAA	---ALL	---TYL	-GLFLGTDY	----	EN--FAAAAAANAWLGLEINMAQI
Sp33	-----PTIL	---	NIA	-GLHMETDI	-N-FS	-LAWF--QAWGGLEINKQAIL
Sp6	-----ASGAI	---	LTL	-GIYLFITLC	-AVIS	-VSWY----LAWLGLEINMMII

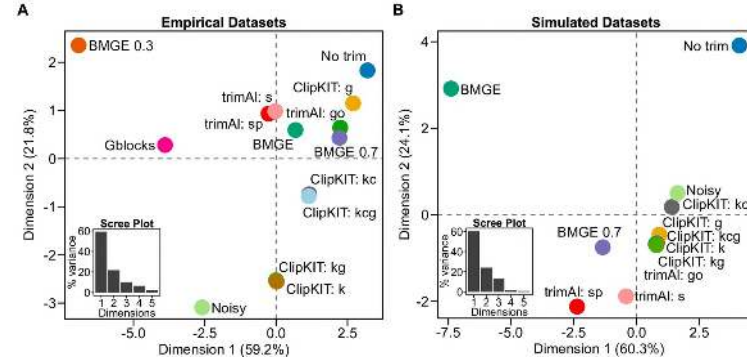
ClipKIT

PLOS BIOLOGY

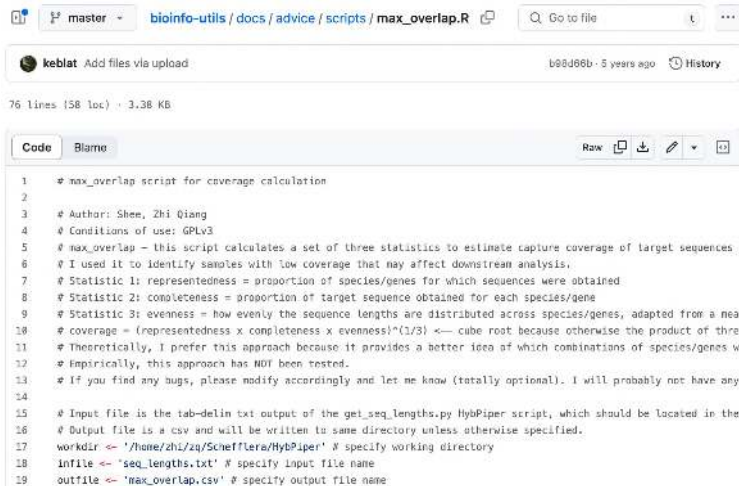
METHODS AND RESOURCES

ClipKIT: A multiple sequence alignment trimming software for accurate phylogenomic inference

Jacob L. Steenwyk^{1*}, Thomas J. Bulda, III², Yuanning Li¹, Xing-Xing Shen³,
Antonios Rokas^{1*}

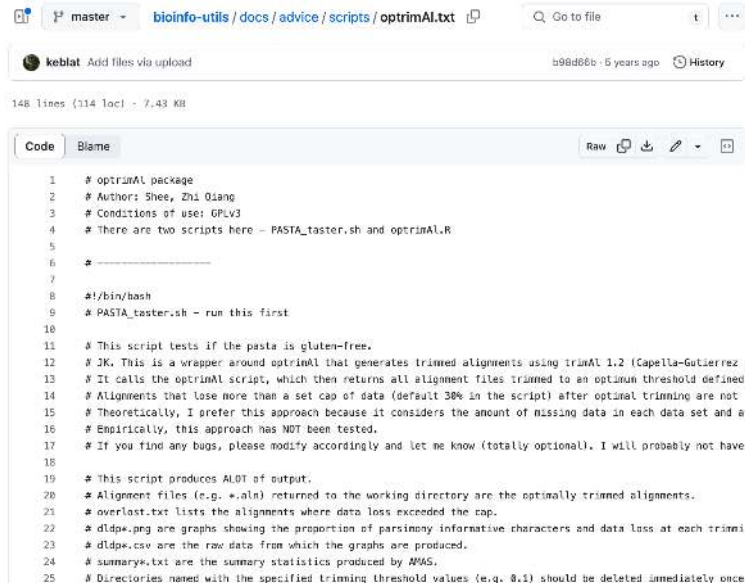


max_overlap



```
1 # max_overlap script for coverage calculation
2
3 # Author: Shee, Zhi Qiang
4 # Conditions of use: GPLv3
5 # max_overlap - this script calculates a set of three statistics to estimate capture coverage of target sequences
6 # I used it to identify samples with low coverage that may affect downstream analysis.
7 # Statistic 1: representedness = proportion of species/genes for which sequences were obtained
8 # Statistic 2: completeness = proportion of target sequence obtained for each species/gene
9 # Statistic 3: evenness = how evenly the sequence lengths are distributed across species/genes, adapted from a near
10 # coverage = (representedness x completeness x evenness)^(1/3) <-- cube root because otherwise the product of three
11 # Theoretically, I prefer this approach because it provides a better idea of which combinations of species/genes w
12 # Empirically, this approach has NOT been tested.
13 # If you find any bugs, please modify accordingly and let me know (totally optional). I will probably not have any
14
15 # Input file is the tab-delim txt output of the get_seq_lengths.py HybPiper script, which should be located in the
16 # Output file is a csv and will be written to same directory unless otherwise specified.
17 workdir <- '/home/zhi/qz/Schnefflera/HybPiper' # specify working directory
18 infile <- 'seq_lengths.txt' # specify input file name
19 outfile <- 'max_overlap.csv' # specify output file name
```

optrimAl



```
1 # optrimAl package
2 # Author: Shee, Zhi Qiang
3 # Conditions of use: GPLv3
4 # There are two scripts here - PASTA_taster.sh and optrimAl.R
5
6 # -----
7
8 #!/bin/bash
9 # PASTA_taster.sh - run this first
10
11 # This script tests if the pasta is gluten-free.
12 # JK. This is a wrapper around optrimAl that generates trimmed alignments using trimAl 1.2 (Capella-Gutierrez
13 # It calls the optrimAl script, which then returns all alignment files trimmed to an optimum threshold defined
14 # Alignments that lose more than a set cap of data (default 30% in the script) after optimal trimming are not
15 # Theoretically, I prefer this approach because it considers the amount of missing data in each data set and a
16 # Empirically, this approach has NOT been tested.
17 # If you find any bugs, please modify accordingly and let me know (totally optional). I will probably not have
18
19 # This script produces ALOT of output.
20 # Alignment files (e.g. *.aln) returned to the working directory are the optimally trimmed alignments.
21 # overlap.txt lists the alignments where data loss exceeded the cap.
22 # dldpe.png are graphs showing the proportion of parsimony informative characters and data loss at each trimm
23 # dldpe.csv are the raw data from which the graphs are produced.
24 # summaryx.txt are the summary statistics produced by APOGAS.
25 # Directories named with the specified trimming threshold values (e.g. 0.1) should be deleted immediately once
```

TreeShrink

Mai and Mirarab *BMC Genomics* 2018, **19**(Suppl 5):272
https://doi.org/10.1186/s12864-018-4620-2

BMC Genomics

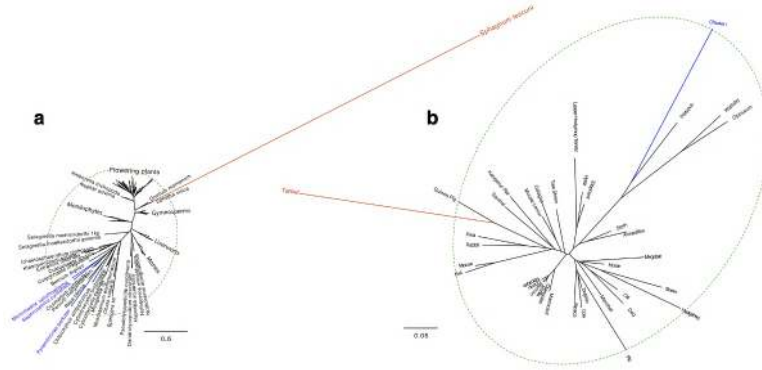
RESEARCH

Open Access

TreeShrink: fast and accurate detection of outlier long branches in collections of phylogenetic trees

Uyen Mai¹ and Siavash Mirarab^{2*}

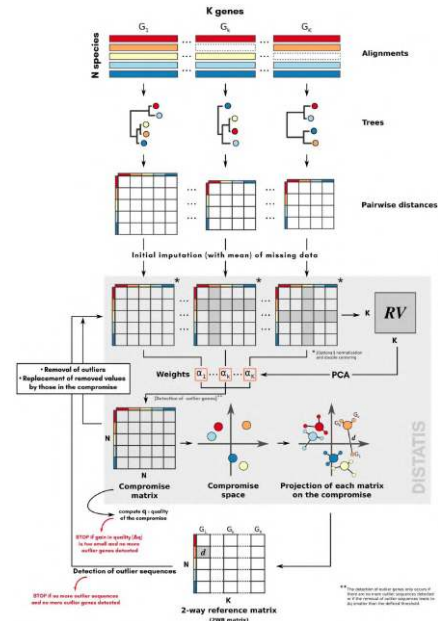
From RECOMB-CG - 2017: The Fifteenth RECOMB Comparative Genomics Satellite Conference
Barcelona, Spain. 04-06 October 2017



PhylteR

PhylteR: Efficient Identification of Outlier Sequences in Phylogenomic Datasets

Aurore Comte,^{1,2,†} Théo Tricou,^{3,†} Eric Tannier,^{3,4} Julien Joseph,^{3,4} Aurélie Siberchicot,³ Simon Penel,³ Rémi Allio,⁵ Frédéric Delsuc,⁶ Stéphane Dray,³ and Damien M. de Vienne^{3,*}



Dry Lab Pipeline

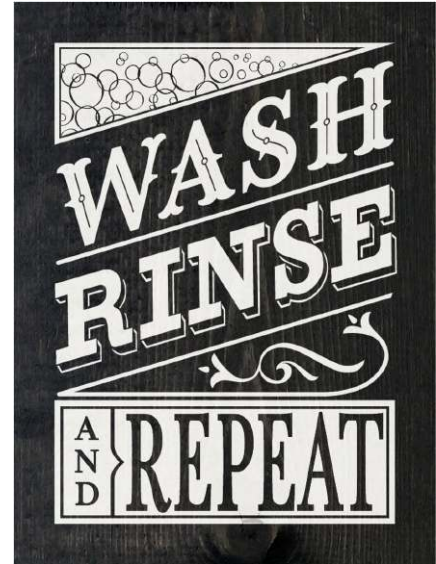
Target
Sequences

MAFFT | MUSCLE | PASTA | UPP

AMAS | PhyKIT | trimAl | ClipKIT

TreeShrink | PhylteR

Aligned
Matrices



Dry Lab Pipeline

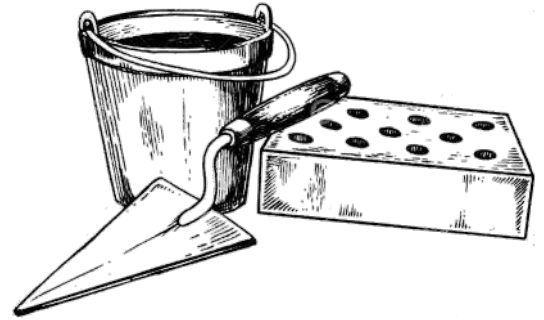
Aligned
Matrices

Brick

Mortar

Masonry

Species
Trees



Dry Lab Pipeline

**Aligned
Matrices**

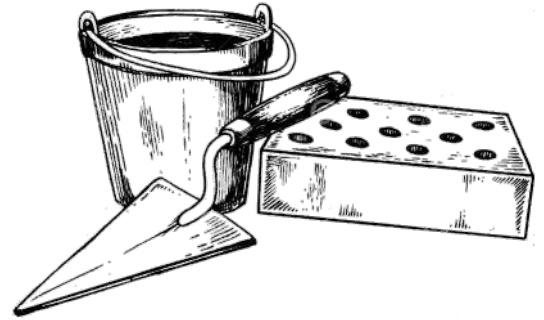
FastTree | IQ-TREE2 | RAxML-NG | PhyML

**Gene
Trees**

Newick utils | Gotree/Goalign

ASTRAL III & PRO | TREE-QMC | treespace

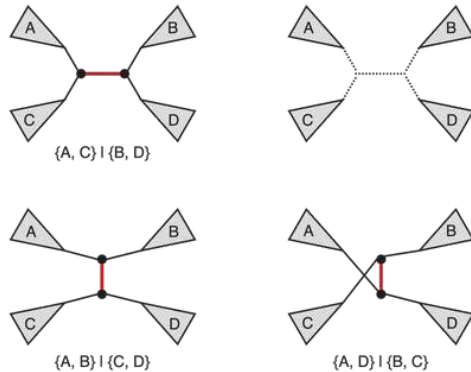
**Species
Trees**



IQ-TREE2

IQ-TREE 2: New Models and Efficient Methods for Phylogenetic Inference in the Genomic Era

Bui Quang Minh ^{*,1,2} Heiko A. Schmidt,³ Olga Chernomor,³ Dominik Schrempf,^{3,4} Michael D. Woodhams,⁵ Arndt von Haeseler,^{1,3,6} and Robert Lanfear^{1,2}



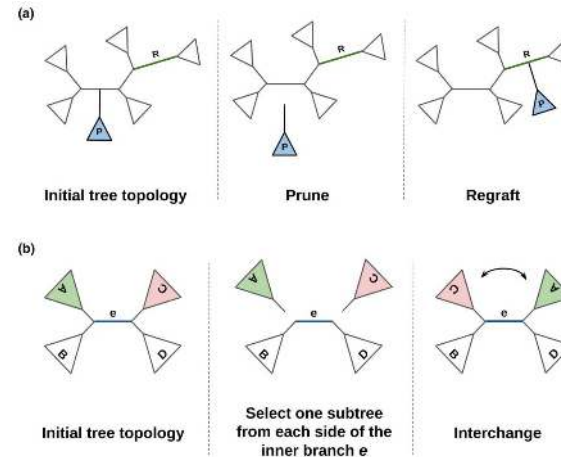
RAxML-NG

Bioinformatics, 2019, 1–3
doi: 10.1093/bioinformatics/btz305
Advance Access Publication Date: 9 May 2019
Application Note

Phylogenetics

RAxML-NG: a fast, scalable and user-friendly tool for maximum likelihood phylogenetic inference

Alexey M. Kozlov ^{1,*}, Diego Darriba ¹, Tomás Flouri¹, Benoit Morel¹ and Alexandros Stamatakis^{1,2}

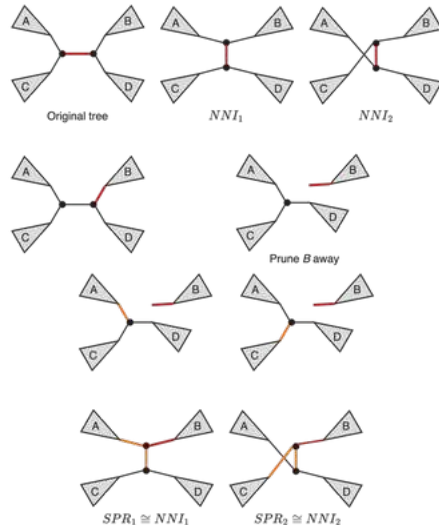


PhyML

Syst. Biol. 59(3):307–321, 2010
 © The Author(s) 2010. Published by Oxford University Press, on behalf of the Society of Systematic Biologists. All rights reserved.
 For Permissions, please email: journals.permissions@oxfordjournals.org
 DOI:10.1093/sysbio/syq010
 Advance Access publication on March 29, 2010

New Algorithms and Methods to Estimate Maximum-Likelihood Phylogenies: Assessing the Performance of PhyML 3.0

STÉPHANE GUINDON^{1,2}, JEAN-FRANÇOIS DUFAYARD¹, VINCENT LEFORT¹, MARIA ANISIMOVA^{1,3,4},
 WIM HORDIJK^{1,5}, AND OLIVIER GASCUEL^{1,*}



FastTree2

OPEN ACCESS Freely available online



FastTree 2 – Approximately Maximum-Likelihood Trees for Large Alignments

Morgan N. Price^{1,2*}, Paramvir S. Dehal^{1,2}, Adam P. Arkin^{1,2,3}

	250	1,250	5,000	78,132
Method	a.a.	a.a.	a.a.	nt.
RAXML 7 (JTT+CAT, SPRs)	90.5%	88.4%	88.4%	–
PhyML 3.0 (JTT+Γ ₄ , SPRs)	89.9%	–	–	–
FastTree 2.0.0 (JTT+CAT or JC+CAT)	86.9%	83.7%	84.3%	92.1%
PhyML 3.0 (JTT+Γ ₄ , no SPRs)	86.0%	–	–	–
FastME 2.06 (log-corrected distances, SPRs)	80.5%	78.8%	77.0%	–
FastTree 2.0.0, no ML NNIs	80.4%	78.3%	76.6%	91.4%
BIONJ (ML distances)	77.7%	73.7%	73.1%	–
Parsimony (RAXML)	76.8%	76.5%	69.4%	–
Neighbor joining (log-corrected distances)	76.0%	72.6%	71.6%	66.1%
Clearcut (log-corrected distances)	75.5%	72.3%	71.5%	58.1%

Newick Utils

BIOINFORMATICS APPLICATIONS NOTE

Vol. 26 no. 13 2010, pages 1669–1670
doi:10.1093/bioinformatics/btq243

Phylogenetics

Advance Access publication May 13, 2010

The Newick utilities: high-throughput phylogenetic tree processing in the UNIX shell

Thomas Junier^{1,2,*} and Evgeny M. Zdobnov^{1,2,3}

Table 1. Selected Newick utilities programs and their functions

Program	Function
nw_clade	Extracts clades (subtrees), specified by labels
nw_distance	Extracts branch lengths in various ways (from root, from parent, as matrix, etc.)
nw_display	Draws trees as ASCII or SVG (suitable for further editing for presentations or publications), several options
nw_match	Reports matches of a tree in a larger tree
nw_order	Orders tree nodes, without altering topology
nw_rename	Changes node labels
nw_reroot	Reroots trees on an outgroup, specified by labels
nw_trim	Trims a tree at a specified depth
nw_topology	Retains topological information

Gotree/Goalign

Published online 11 August 2021

NAR Genomics and Bioinformatics, 2021, Vol. 3, No. 3 1
<https://doi.org/10.1093/nargab/lqab075>

Gotree/Goalign: toolkit and Go API to facilitate the development of phylogenetic workflows

Frédéric Lemoine^{1,2,*} and Olivier Gascuel¹

Table 1. Subset of commands implemented in Gotree/Goalign

Goalign Command	Description
reformat	Converts between Nexus, Clustal, Fasta and Phylip
rename	Modifies or cleans sequence names
clean	Removes sites/sequences
codonalign	Aligns by codons using a protein alignment
concat	Concatenates several alignments
mask	Masks parts of the alignment
translate	Translates input alignment in amino-acids
extract	Extracts subsequences from input alignments
build seqboot	Builds bootstrap alignments
compute distance	Computes distance matrix
stats	Computes statistics about input alignments
Gotree Command	Description
reformat	Converts between Newick, Nexus and PhyloXML
prune	Removes user-defined tips
collapse	Collapses branches
reroot	Reroots trees
compare trees	Compares tree topologies (bipartitions)
compute support	Computes branch supports (30) and (31)
stats	Computes statistics about input trees
matrix	Computes patristic distance matrix
upload itol	Uploads a tree to iTOL (9)

ASTRAL-III

Zhang et al. *BMC Bioinformatics* 2018, **19**(Suppl 6):153
<https://doi.org/10.1186/s12859-018-2129-y>

BMC Bioinformatics

RESEARCH

Open Access



ASTRAL-III: polynomial time species tree reconstruction from partially resolved gene trees

Chao Zhang³, Maryam Rabiee², Erfan Sayyari¹ and Siavash Mirarab^{1*}

ASTRAL-Pro

ASTRAL-Pro: Quartet-Based Species-Tree Inference despite Paralogy

Chao Zhang,¹ Celine Scornavacca,² Erin K Molloy,³ and Siavash Mirarab ^{*,4}

Accurate Species Tree Estimator (ASTER*)



A family of optimization algorithms for species tree inference:

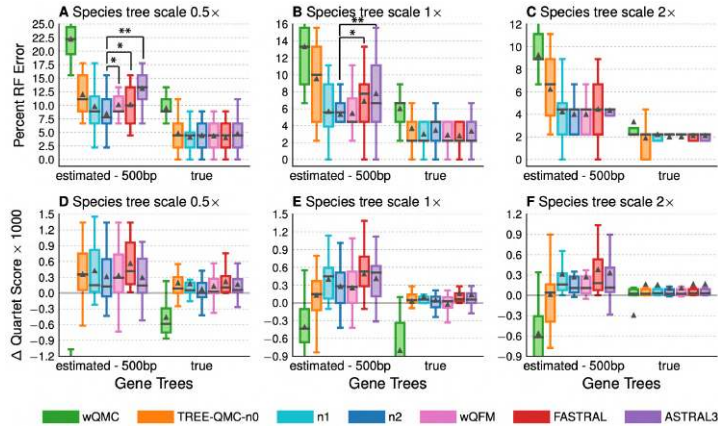
1. [ASTRAL](#)
2. [ASTRAL-Pro](#)
3. [Weighted ASTRAL](#)
4. [CASTER-site](#)
5. [CASTER-pair](#)
6. [WASTER-site](#)
7. SISTER
8. MONSTER

TREE-QMC

Improving quartet graph construction for scalable and accurate species tree estimation from gene trees

Running Title: TREE-QMC

Yunheng Han^{1,2} and Erin K. Molloy^{1,2,*}



treespace

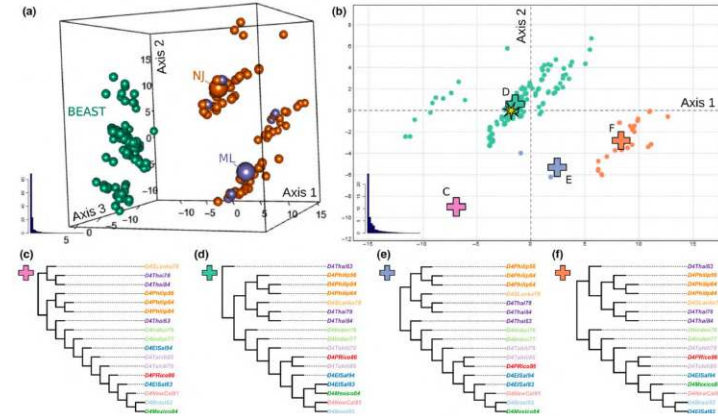
Received: 22 November 2016 | Revised: 17 March 2017 | Accepted: 21 March 2017
DOI: 10.1111/1755-0998.12676

RESOURCE ARTICLE

WILEY **MOLECULAR ECOLOGY**
RESOURCES

TREESPACE: Statistical exploration of landscapes of phylogenetic trees

Thibaut Jombart^{1,*} | Michelle Kendall^{2,*} | Jacob Almagro-Garcia³ | Caroline Colijn²



DiscoVista

Molecular Phylogenetics and Evolution 122 (2018) 110–115

Contents lists available at ScienceDirect



Molecular Phylogenetics and Evolution

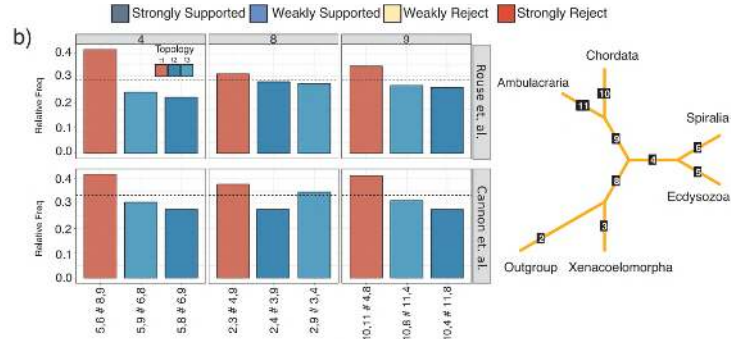
journal homepage: www.elsevier.com/locate/ympev



Short Communication

DiscoVista: Interpretable visualizations of gene tree discordance

Erfan Sayyari^a, James B. Whitfield^b, Siavash Mirarab^{a,*}

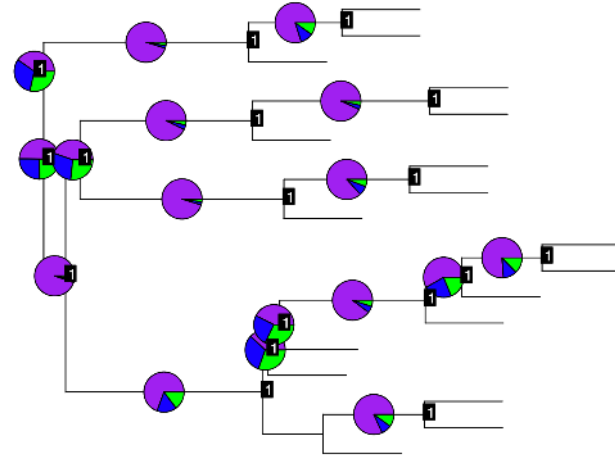


AstralPlane

AstralPlane

R Package For Preparing, Running, Analyzing and Plotting from the Species Tree Program ASTRAL-III

This R package is meant to facilitate ASTRAL-III analyses and provide easy R plotting. The package helps prepare analyses from a folder of gene trees, runs astral from R, and creates a new S4 object type "AstralPlane" for easily analyzing the output from ASTRAL-III. The package provides several different types of plots, from pie charts on phylogenetic trees representing the quartet frequencies to plotting the gene tree frequencies as far plots.



TreePL

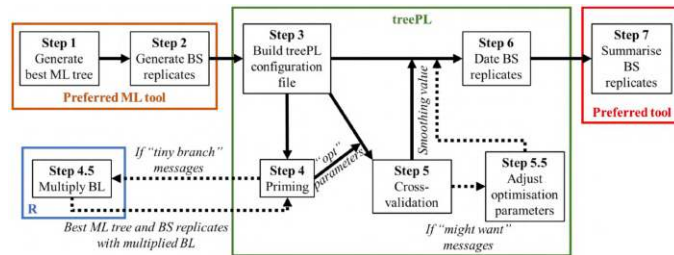
BIOINFORMATICS APPLICATIONS NOTE Vol. 28 no. 20 2012, pages 2689–2690
doi:10.1093/bioinformatics/bts492

Phylogenetics

Advance Access publication August 20, 2012

treePL: divergence time estimation using penalized likelihood for large phylogenies

Stephen A. Smith^{1,*} and Brian C. O'Meara²



Copyright Kevin J. L. Maurin, Creative Commons Attribution License (CC BY 4.0).

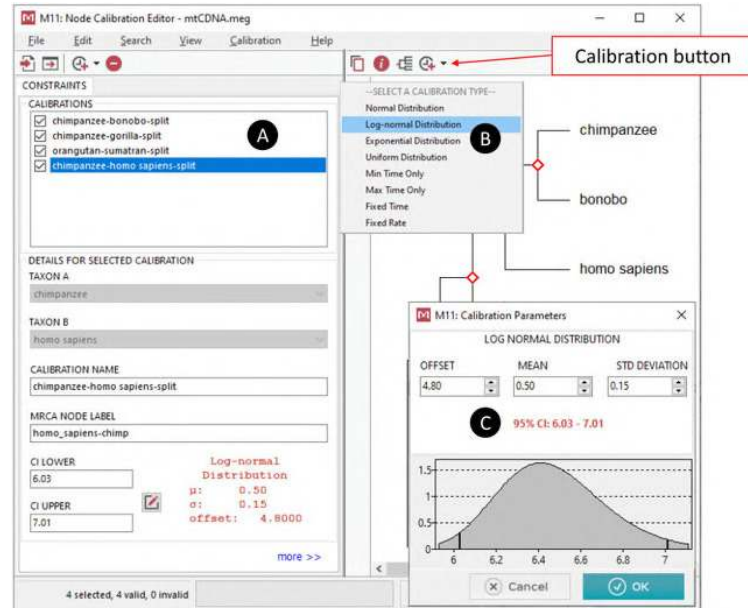
An empirical guide for producing a dated phylogeny with treePL in a maximum likelihood framework

Kévin J. L. Maurin

RelTime

Theoretical Foundation of the RelTime Method for Estimating Divergence Times from Variable Evolutionary Rates

Koichiro Tamura,^{1,2} Qiqing Tao,^{3,4} and Sudhir Kumar^{*,3,4,5}



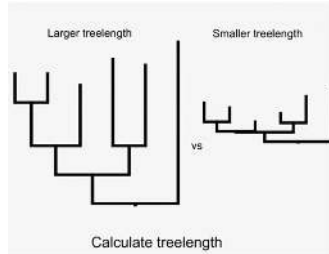
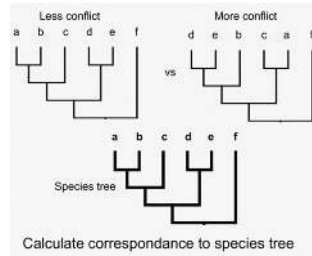
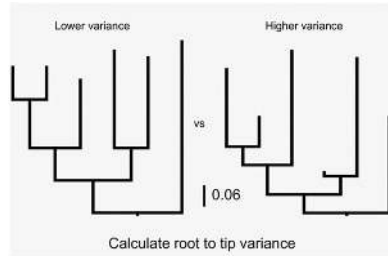
SortaDate



RESEARCH ARTICLE

So many genes, so little time: A practical approach to divergence-time estimation in the genomic era

Stephen A. Smith^{1*}, Joseph W. Brown^{2,3}, Joseph F. Walker^{1,4}



genesortR

Phylogenomic Subsampling and the Search for Phylogenetically Reliable Loci

Nicolás Mongiardino Koch*

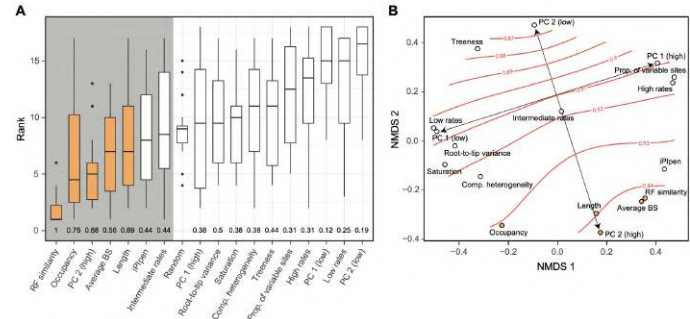
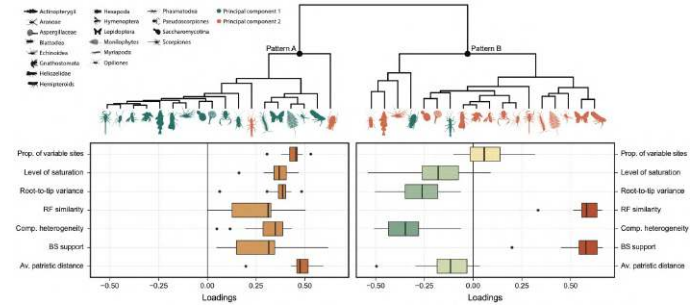


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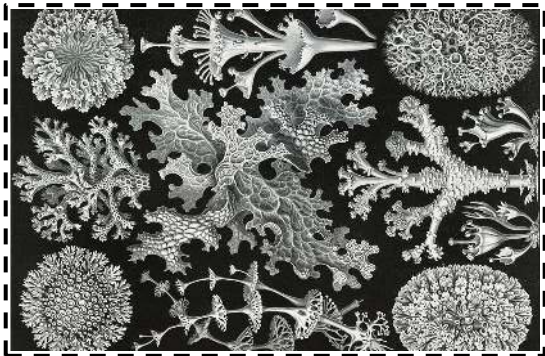
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Natural History Collections



Biodiversity Genomics

Natural History Collections

04

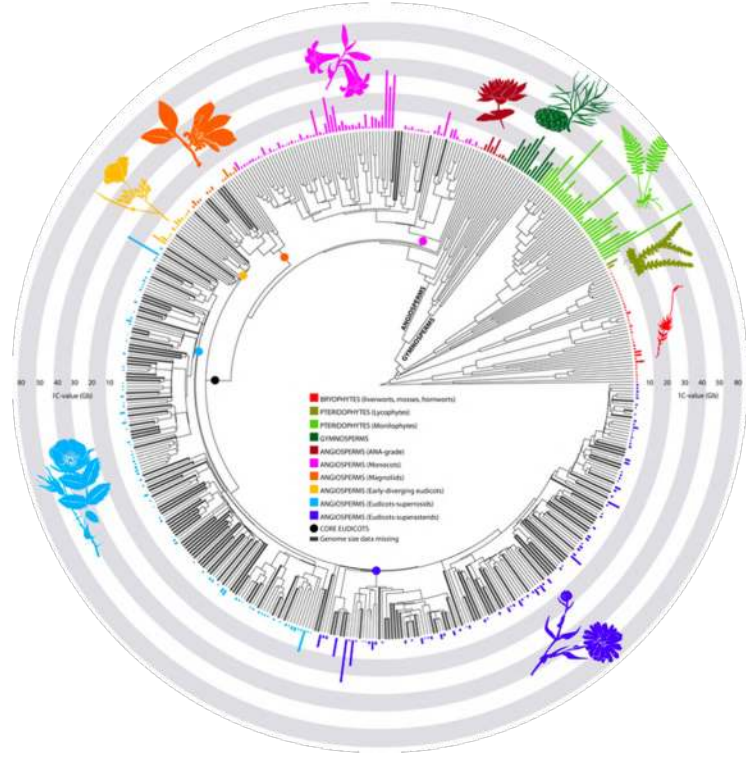
Explaining the fate of diversity

Understanding the
eco-evolutionary
processes that have
shaped the traits
leading to its origin
and maintenance



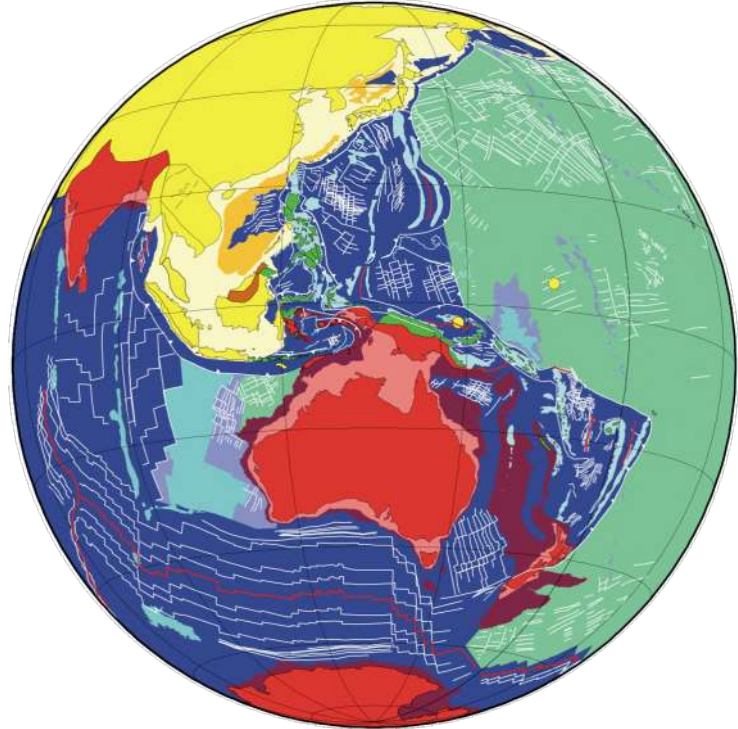
Understanding eco-evolutionary processes

Taking into account
intrinsic factors...



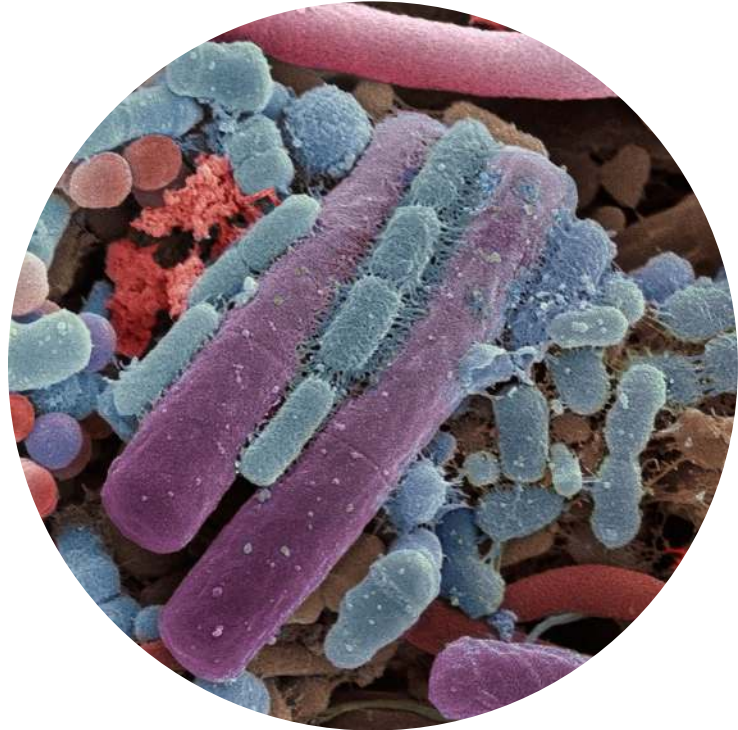
Understanding eco-evolutionary processes

...and extrinsic factors
that, in turn, can
be abiotic...



Understanding eco-evolutionary processes

...or biotic



Understanding eco-evolutionary processes

The information
stored in
genomes is a
record of past
events...



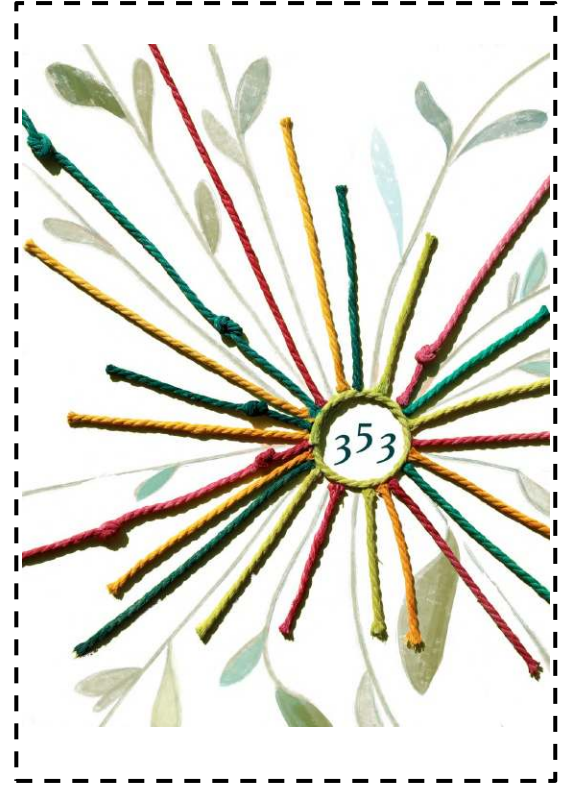
Understanding eco-evolutionary processes

...also registered
in the fossil
record



Angiosperms353

A universal tool for flowering plant systematics



Angiosperms353

Classification: flowering plants, ~ 370K spp. (APG VI: 416 fams. & 64 orders)

Importance: agronomic, edible, timber, pulp, fiber, meds, ornamental, fuel, etc.

Available resources:

655 transcriptomes  **1KP**

20 genomes 

Hamming dist., k-medoids clustering

Daicel Arbor Biosciences 

Resulting kit:

353 nuclear SCOGs

~260,8 Kbp matrix

~75,2K 120-mer 3x tiled probes

Syst. Biol. 68(4):594–606, 2019

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DOI:10.1093/sysbio/syy186

Advance Access publication December 10, 2018

A Universal Probe Set for Targeted Sequencing of 353 Nuclear Genes from Any Flowering Plant Designed Using k-Medoids Clustering

MATTHEW G. JOHNSON^{1,2,*}, LISA POKORNY³, STEVEN DODSWORTH^{3,4}, LAURA R. BOTIGUE^{3,5}, ROBYN S. COWAN³, ALISON DEVAULT⁶, WOLF L. EISERHARDT^{3,7}, NIROSHINI EPITAWALAGE³, FELIX FOREST³, JAN T. KIM³, JAMES H. LEEBENS-MACK⁸, ILIA J. LEITCH³, OLIVIER MAURIN³, DOUGLAS E. SOLTIS^{9,10}, PAMELA S. SOLTIS^{9,10}, GANE KA-SHU WONG^{11,12,13}, WILLIAM J. BAKER³, AND NORMAN J. WICKETT^{2,14}

¹Department of Biological Sciences, Texas Tech University, Lubbock, TX 79409, USA; ²Plant Science and Conservation, Chicago Botanic Garden, 1000 Lake Cook Road, Glenview, IL 60022, USA; ³Department of Comparative Plant and Fungal Biology, Royal Botanic Gardens, Kew, Richmond, Surrey TW9 3AE, UK; ⁴School of Life Sciences, University of Bedfordshire, University Square, Luton LU1 3JU, UK; ⁵Centre for Research in Agricultural Genomics, Campus UAB, Edifici CRAG, Bellaterra Cerdanyola del Vallès, 08193 Barcelona, Spain; ⁶Arbor Biosciences, 5840 Interface Dr, Suite 101, Ann Arbor, MI 48103, USA; ⁷Department of Bioscience, Aarhus University, 8000 Aarhus C, Denmark; ⁸Department of Plant Biology, University of Georgia, 2502 Miller Plant Sciences, Athens, GA 30602, USA; ⁹Department of Biology, University of Florida, 220 Bartram Hall, Gainesville, FL 32611-8525, USA; ¹⁰Florida Museum of Natural History, University of Florida, 3215 Hull Road, Gainesville, FL 32611-2710, USA; ¹¹BGI-Shenzhen, Beishan Industrial Zone, Yantian District, Shenzhen 518083, China; ¹²Department of Biological Sciences, University of Alberta, Edmonton, AB T6G 2E9, Canada; ¹³Department of Medicine, University of Alberta, Edmonton, AB T6G 2E1, Canada; and ¹⁴Program in Plant Biology and Conservation, Northwestern University, 2205 Tech Drive, Evanston, IL 60208, USA

*Correspondence to be sent to: Department of Biological Sciences, Texas Tech University, Lubbock, TX 79409, USA;

E-mail: matt.johnson@ttu.edu.

Matthew G. Johnson, Lisa Pokorny, Steven Dodsworth contributed equally to this article.

Flowering plants (Angiospermae)

APG VI classification (2016): ~370K species, ~13.6 genera, 416 families, & 64 orders
Importance: 90% terrestrial spp., nutrition, fuels, lumber, meds, cultural, etc.

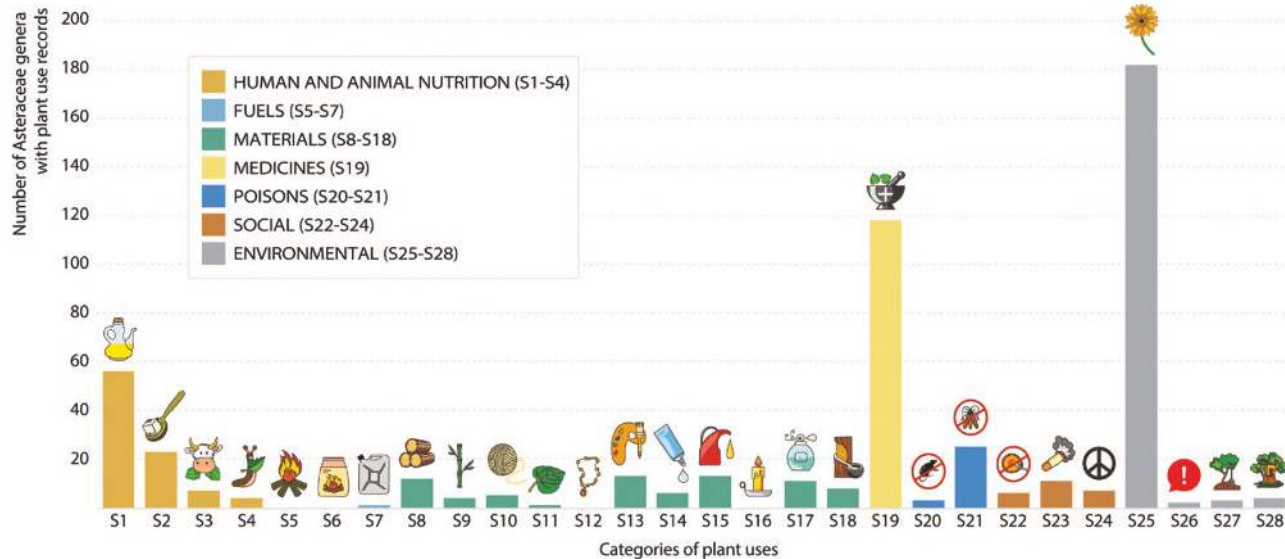
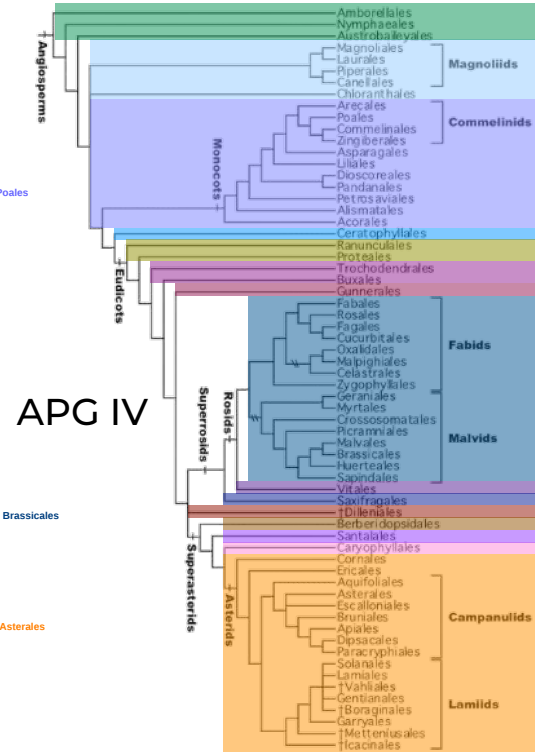
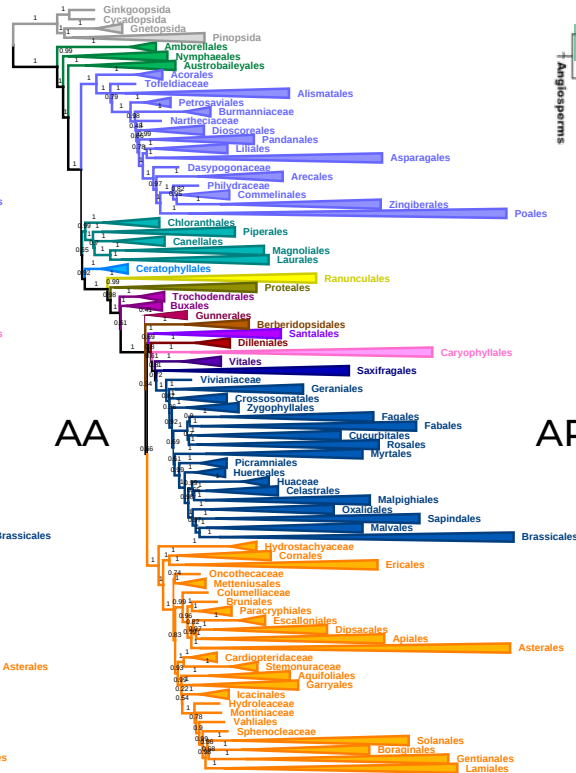
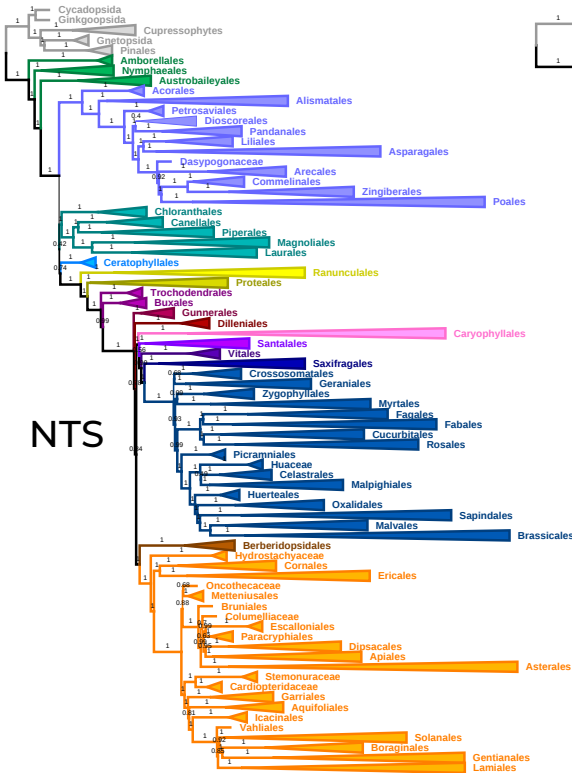


Fig. 2. Palazzesi et al. (2022). *Bot. J. Linn. Soc.* 200(2): 143–164.

Angiosperms353



Angiosperms353



Commelinales: ~885 spp. (70 gen., 5 fams.)

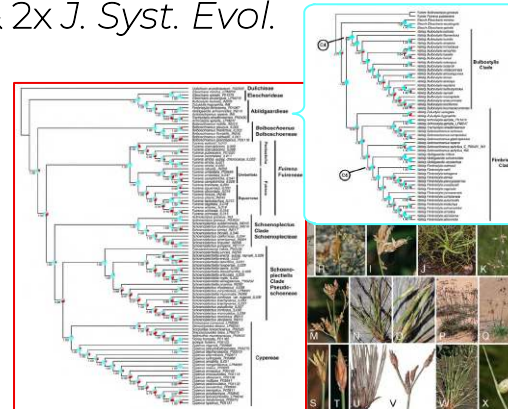
Interest: ornamental (kangaroo paws, spiderworts), invasive (water hyacinth)

Publications: *Am. J. Bot.*

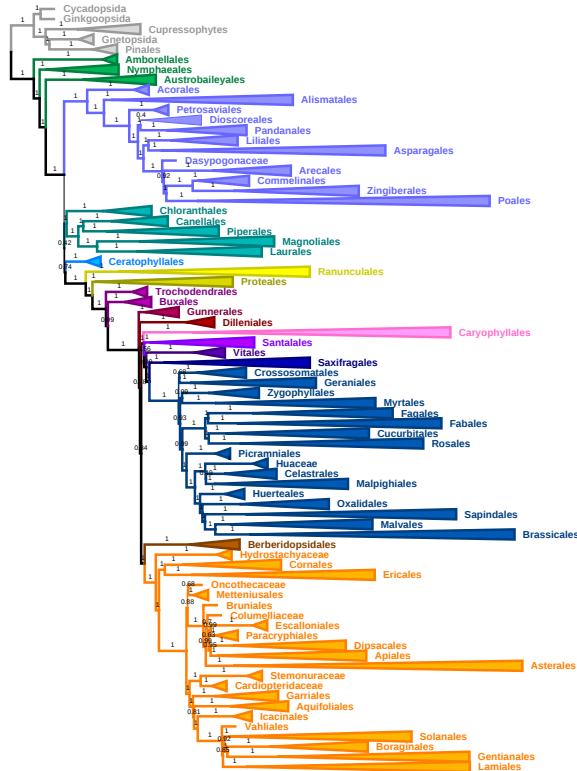
- **Cyperaceae** (Poales): ~5,6K spp. (~90 gen.)

Interest: edible (chufa), pulp (papyrus)

Publications: *Front. Plant Sci.*, *Bot. J. Linn. Soc.*, & 2x *J. Syst. Evol.* 



Angiosperms353



frontiers
in Plant Science

OPEN ACCESS
EDITED BY: JAMES H. WILSON,
University of Oxford, United Kingdom



Tackling Rapid Radiations With Targeted Sequencing

Isabel Larridon^{1,2,3,4}, Tamara Villaverde^{5,6,7,8,9,10}, Alexandre R. Zuntini^{1,2,3}, Lisa Pokorny^{1,2,3}, Grace E. Brewer^{1,2,3}, Niroshini Eptawalage¹, Isabel Fairlie^{1,2}, Etienne Lévêillé-Bourret^{1,2}, Jean Kim¹, Enrique Obispo^{1,2}, Olivier Mauch¹, Martin Kott¹, Andrew L. Page¹, Felix Forest¹ and William J. Baker¹

Botanical Journal of the Linnean Society, 2021, XX, 1–26. With 6 figures.

Resolving generic limits in Cyperaceae tribe Abildgaardieae using targeted sequencing

ISABEL LARRIDON^{1,2,3,4}, ALEXANDRE R. ZUNTINI^{1,2,3}, RUSSELL L. BARRETT¹, KAREN L. WILSON¹, JEREMY J. BRUHL¹, PAUL GOETGHEBEUR¹, WILLIAM J. BAKER¹, GRACE E. BREWER¹, NIROSHINI EPTAWALAGE¹, ISABEL FAIRLIE^{1,2}, FELIX FOREST¹, IZAI A. B. SABINO KIKUCHI^{1,2}, LISA POKORNY^{1,2,3}, ILIAS SEMMOURI¹, DANIEL SPALINK¹, DAVID A. SIMPSON¹, A. MUTHAMA MUASYA^{1,2,3} and ERIC H. ROALSON^{1,2}

Botany



INVITED SPECIAL ARTICLE
For the special issue: Tackling rapid radiations in plants: tools for resolving their phylogenies

A comprehensive phylogenomic study of the monocot order Commelinales, with a new classification of Commelinaceae

Alexandre R. Zuntini^{1,2,3}, James H. Wilson^{1,2}, Lisa Pokorny^{1,2,3}, Felix Forest^{1,2}, and William J. Baker^{1,2}

JSE Journal of Systematics and Evolution

doi: 10.1111/jse.12721

Research Article

Targeted sequencing supports morphology and embryo features in resolving the classification of Cyperaceae tribe Fuireneae s.l.

Julien R. Stan^{1,2}, Pedro Jiménez-Mejías^{1,3}, Alexandre R. Zuntini^{1,2,3}, Etienne Lévêillé-Bourret^{1,2}, Elias Semmouri¹, Muthama Muasya^{1,2}, William J. Baker¹, Grace E. Brewer¹, Niroshini Eptawalage¹, Isabel Fairlie^{1,2}, Felix Forest¹, Izai A. B. Sabino Kikuchi^{1,2}, Lisa Pokorny^{1,2,3}, and Isabel Larridon^{1,2,3,4}

JSE Journal of Systematics and Evolution

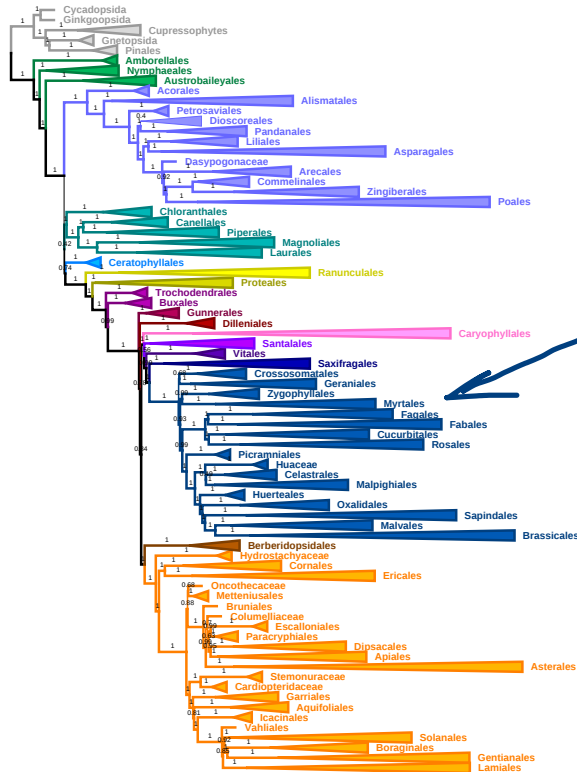
doi: 10.1111/jse.12757

Research Article

A new classification of Cyperaceae (Poales) supported by phylogenomic data

Isabel Larridon^{1,2,3,4}, Alexandre R. Zuntini^{1,2,3}, Etienne Lévêillé-Bourret^{1,2}, Russell L. Barrett¹, Julien R. Stan¹, A. Muthama Muasya^{1,2}, Tamara Villaverde^{5,6,7,8,9,10}, Kenneth Baudens¹, Grace E. Brewer¹, Jeremy J. Bruhl¹, Suzana M. Costa¹, Tammy L. Elliott¹, Niroshini Eptawalage¹, Marcial Escudero¹, Isabel Fairlie^{1,2}, Paul Goetghebeur¹, Andrew L. Page^{1,2,3}, Pedro Jiménez-Mejías^{1,3}, Izai A. B. Sabino Kikuchi^{1,2}, Madesto Lucero¹, José Ignacio Márquez-Correa¹, Santiago Martín-Bravo¹, Olivier Mauch¹, Lisa Pokorny^{1,2,3}, Eric H. Roalson^{1,2}, Elias Semmouri¹, David A. Simpson¹, Daniel Spalink¹, W. Wayt Thomas¹, Karen L. Wilson¹, Martin Zechner¹, Felix Forest¹, and William J. Baker¹

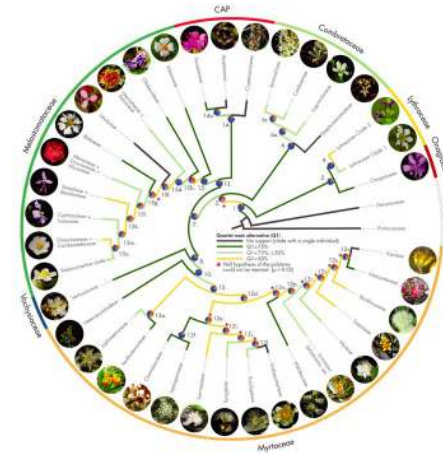
Angiosperms353



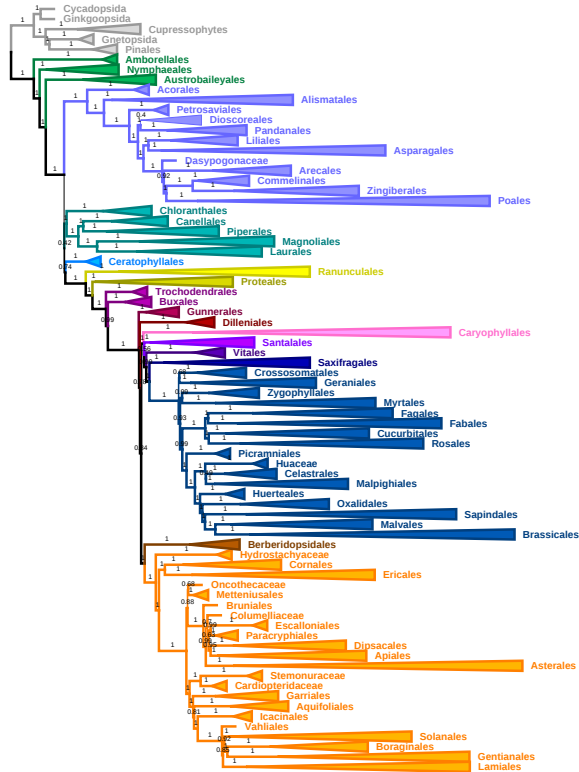
Myrtales: ~13K spp. (~380 gen. 9 fams.)

Interest: edible (fodder), edible (pomegranate, guava, cloves), ornamental (fuchsia, evening primroses), pulp (eucalypts), essential oils (sundrops)

Publications: *Am. J. Bot.*



Angiosperms353



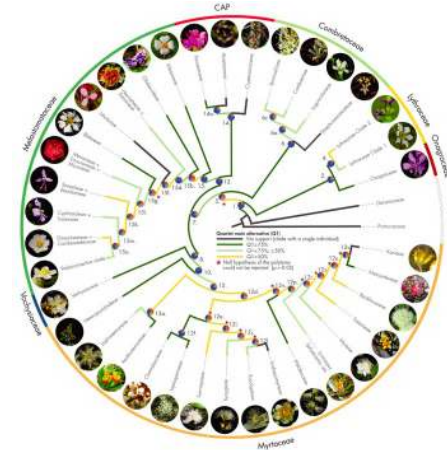
Angiosperm Journal of
Botany



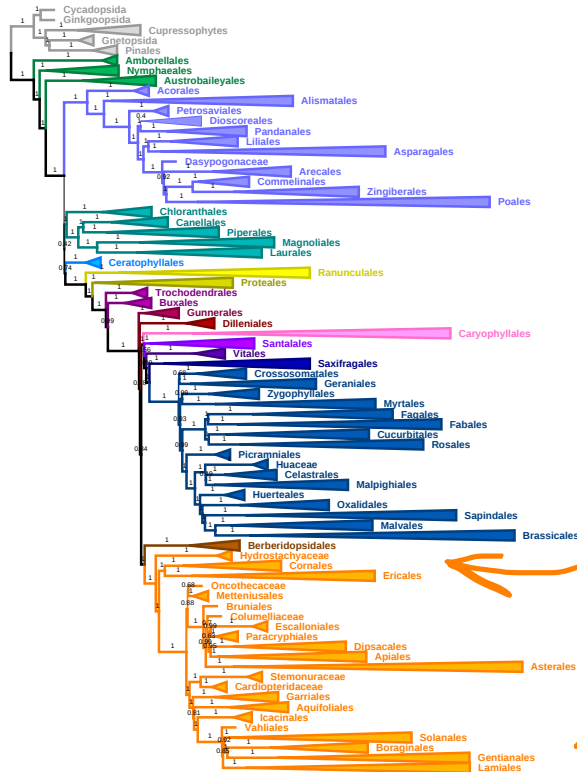
INVITED SPECIAL ARTICLE

For the Special Issue: Exploring Angiosperms353: a Universal Toolkit for Flowering Plant Phylogenomics

A nuclear phylogenomic study of the angiosperm order Myrtales, exploring the potential and limitations of the universal Angiosperms353 probe set



Angiosperms353



Cornales: ~600 spp. (~50 gen., 6 fams.)

Interest: ornamental (hortensias, mock-oranges, dogwoods), lumber (tupelos)

Articles: *Am. J. Bot.*

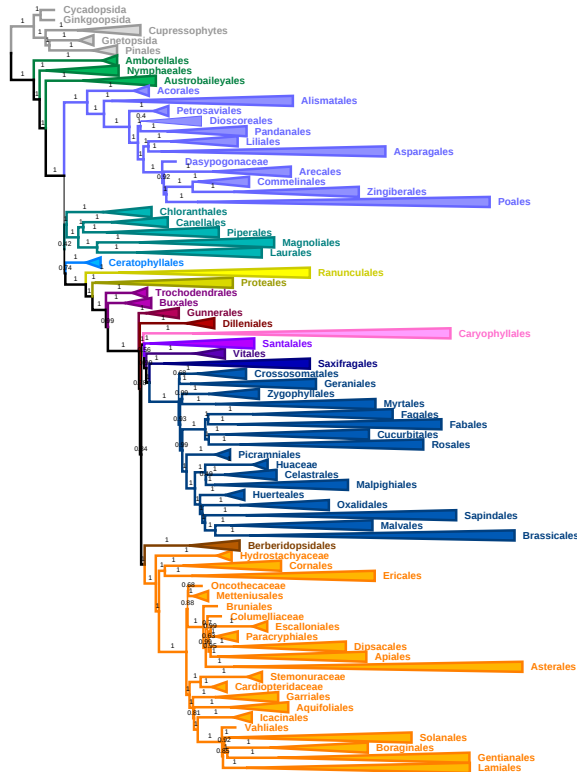
Gentianales: ~22,5K spp. (1,1K gen., 5 fams.)

Interest: drinks (coffee), food (soursop), meds (quinine, strychnine, vincristine), ornamental (gardenia, oleander), dyes.

Articles: *Am. J. Bot.*



Angiosperms353



American Journal of
Botany



RESEARCH ARTICLE

INVITED SPECIAL ARTICLE

For the Special Issue: Exploring Angiosperms353: a Universal Toolkit for Flowering Plant Phylogenomics

Comprehending Coriales: phylogenetic reconstruction of the order using the Angiosperms353 probe set

Shawn K. Thomas^{1,2*}, Xiang Liu^{1*}, Zhi-Yuan Du¹, Yibo Dong^{1*}, Amanda Cummings¹, Lisa Pokorny^{1*}, Qui-Yun (Jenny) Xiang^{1,3}, and James H. Leebens-Mack^{1*}

American Journal of
Botany



RESEARCH ARTICLE

INVITED SPECIAL ARTICLE

For the Special Issue: Exploring Angiosperms353: a Universal Toolkit for Flowering Plant Phylogenomics

Settling a family feud: a high-level phylogenomic framework for the Gentianales based on 353 nuclear genes and partial plastomes

Alexandre Antonelli^{1,2,3,4*}, James J. Clarkson^{1,5}, Kent Kainulainen¹, Olivier Maurin¹, Grace E. Brewer¹, Aaron P. Davis¹, Niroshini Epitawalage¹, David J. Goyder¹, Tatjana Livshultz¹, Claes Persson¹, Lisa Pokorny^{1*}, Shannon C. K. Straub¹, Lena Struwe¹, Alexandre R. Zuntini¹, Felix Forest^{1*}, and William J. Baker^{1*}



Flowering Plant Quipu

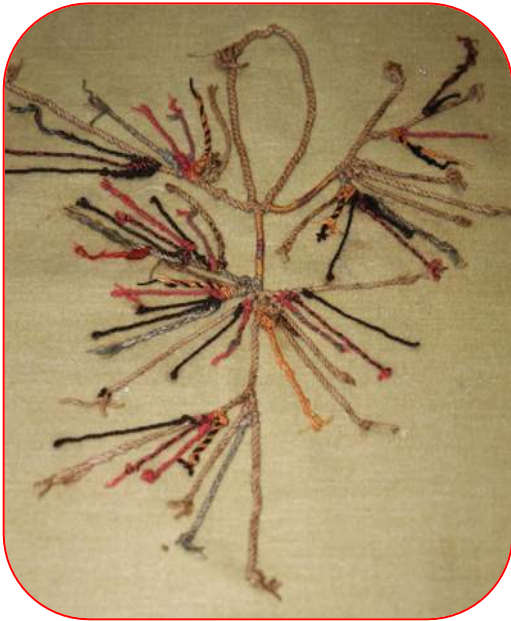
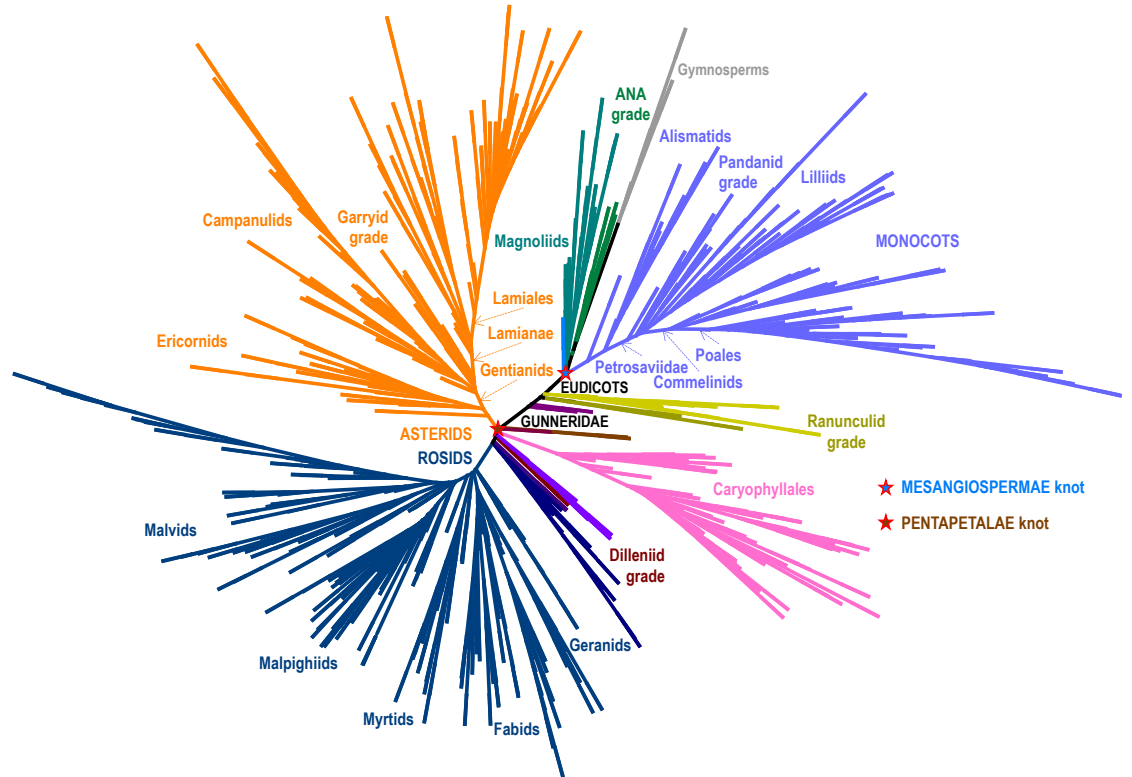


Fig. 8. Loop-and-branch-type khipu. Urton. 2014. *Antiquity*.



Population Genomics



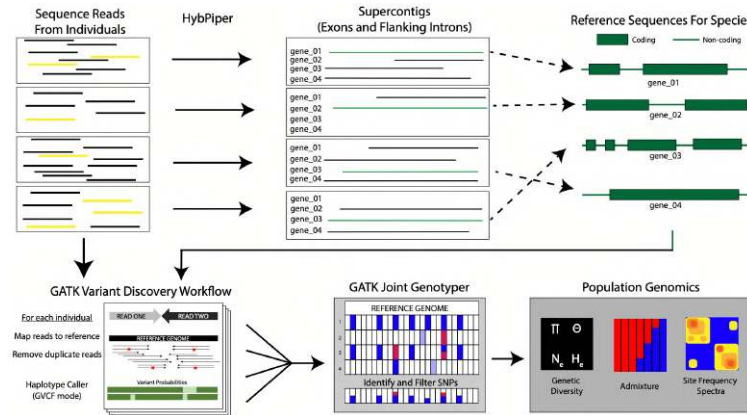
APPLICATION ARTICLE

INVITED SPECIAL ARTICLE

For the Special Issue: Exploring Angiosperms353: A Universal Toolkit for Flowering Plant Phylogenomics

On the potential of Angiosperms353 for population genomic studies

Madeline Slimp¹, Lindsay D. Williams¹, Haley Hale¹, and Matthew G. Johnson^{1,2}



Systematic Botany (2023), 98(4): pp. 1107–1113
© Copyright 2023 by the American Society of Plant Taxonomists
DOI 10.1002/syst.14213.35701080806
Date of publication December 21, 2023

Are Palmer's Elm-Leaf Goldenrod and the Smooth Elm-Leaf Goldenrod Real? The Angiosperms353 Kit Provides Within-Species Signal in *Solidago ulmifolia* s. l.

James B. Beck,^{1,2,5} Morgan L. Markley,¹ Mackenzie G. Zielke,¹ Justin R. Thomas,³ Haley J. Hale,⁴ Lindsay D. Williams⁵, and Matthew G. Johnson¹

¹Department of Biological Sciences, Wichita State University, Wichita, Kansas 67260, USA

²Botanical Research Institute of Texas, Fort Worth, Texas 76107, USA

³NatureCITE, Springfield, Missouri 65803, USA

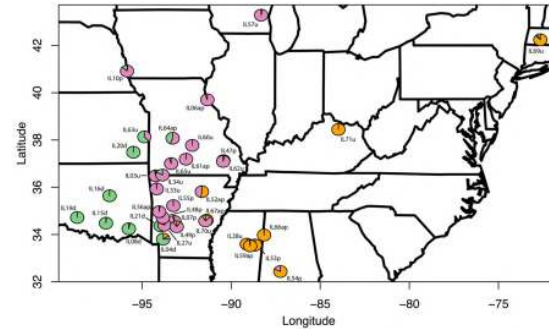
⁴Department of Biological Sciences, Texas Tech University, Lubbock, Texas 79409, USA

⁵Author for correspondence (james.beck@wichita.edu)

Communicating Editor: Ashley B. Morris

Abstract—The genus *Solidago* represents a taxonomically challenging group due to its sheer number of species, putative hybridization, polyploidy, and shallow genetic divergences among species. Here we use a dataset obtained exclusively from herbarium specimens to evaluate the status of *Solidago ulmifolia* var. *palmeri*, a morphologically subtle taxon potentially confined to Alabama, Arkansas, Mississippi, and Missouri. A multivariate analysis of both discrete and continuous morphological data revealed no clear distinction between *S. ulmifolia* var. *palmeri* and *Solidago ulmifolia* var. *ulmifolia*. *Solidago ulmifolia* var. *palmeri*'s status was also assessed with a phylogenomic and SNP clustering analysis of data generated with the “Angiosperms353” probe kit. Neither analysis supported *Solidago ulmifolia* var. *palmeri* as a distinct taxon, and we suggest that this name should be discarded. The status of *Solidago delicatula* (formerly known as *Solidago ulmifolia* var. *microphylla*) was also assessed. Both morphological and phylogenetic analyses supported the species status of *S. delicatula* and we suggest maintaining this species at its current rank. These results highlight the utility of the Angiosperms353 probe kit, both with herbarium tissue and at lower taxonomic levels. Indeed, this is the first study to utilize this kit to identify genetic groups within a species.

Keywords—hyb-seq, North America, Ozark Mountains, *Solidago delicatula*, *S. ulmifolia* var. *palmeri*, species delimitation.



Museomics

Herbariomics / Fungariomics

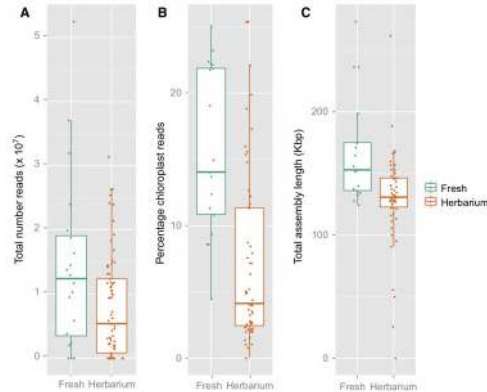
Integrating Natural History Collections into
Biodiversity Research



Herbariomics / Fungariomics

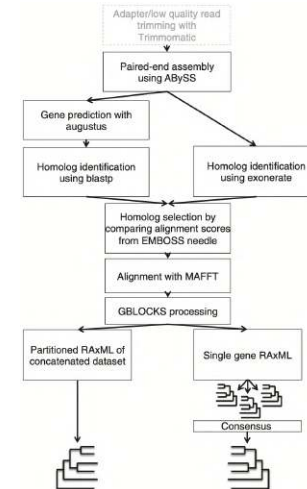
Herbarium genomics: plastome sequence assembly from a range of herbarium specimens using an Iterative Organelle Genome Assembly pipeline

FRECK T. BAKKER¹*, DI LEI¹, JIAYING YU¹, SETAREH MOHAMMADIN¹, ZHEN WU¹, SARA VAN DE KERKE¹, BARBARA GRAVENDEEL^{2,3,4}, MATHILIS NIEUWENHUIS¹, MARTIJN STAATS¹, DAVID E. ALQUEZAR-PLANAS⁵ and RENS HOLMER¹



Tales from the crypt: genome mining from fungarium specimens improves resolution of the mushroom tree of life

BRYN T. M. DENTINGER^{1,2*}, ESTER GAYA¹, HEATH O'BRIEN³, LAURA M. SLIZ², ROBERT LACHLAN⁴, JORGE R. DÍAZ-VALDERRAMA⁵, RACHEL A. KOCH⁶ and M. CATHERINE AIME²



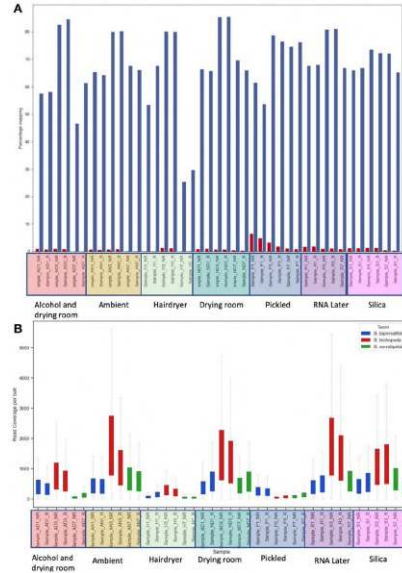
Herbariomics

frontiers
in Ecology and Evolution

ORIGINAL RESEARCH
published: 01 November 2018
doi: 10.3389/fecol.2018.00020

The Limits of Hyb-Seq for Herbarium Specimens: Impact of Preservation Techniques

Laura L. Forrest¹, Michelle L. Hart¹, Mark Hughes¹, Hannah R. Wilson^{1,2},
Kue-Fang Chung¹, Yu-Hsin Tseng¹ and Catherine A. Kitchin^{1,2*}

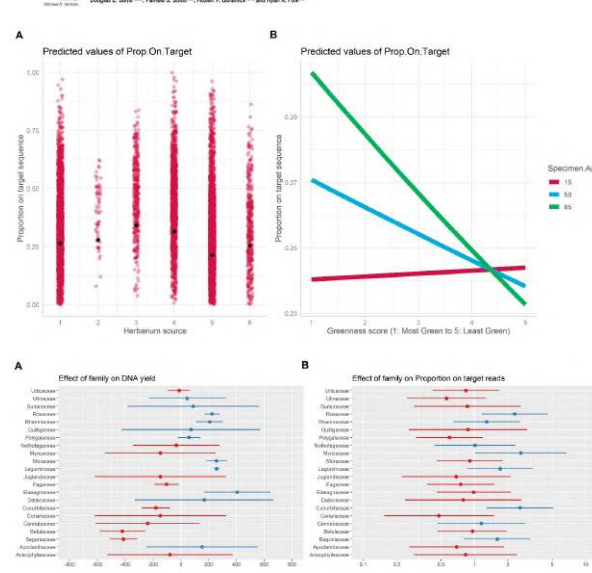


frontiers
in Plant Science

ORIGINAL RESEARCH
published: 01 November 2018
doi: 10.3389/fpls.2018.00020

The Effects of Herbarium Specimen Characteristics on Short-Read NGS Sequencing Success in Nearly 8000 Specimens: Old, Degraded Samples Have Lower DNA Yields but Consistent Sequencing Success

Heather R. Kuhn¹, Joshua R. Davis¹, Carol M. Steinhilber¹, Nathaniel LaFrance¹,
Douglas C. Soltis¹, Pamela S. Soltis¹, Robert R. Goodrich^{1,2} and Ryan A. Puck^{1,2*}

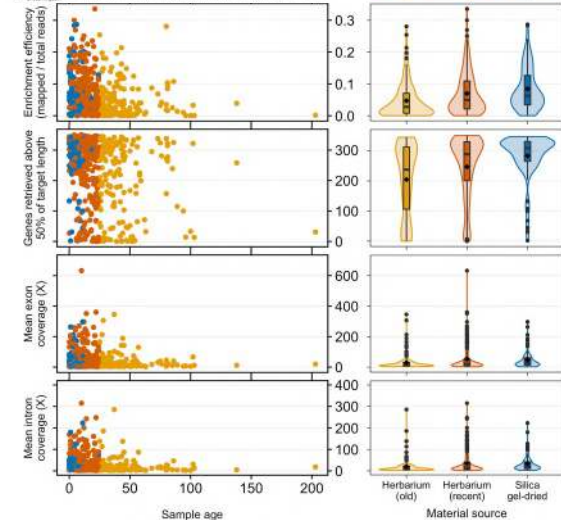


frontiers
in Plant Science

ORIGINAL RESEARCH
published: 01 November 2018
doi: 10.3389/fpls.2018.00020

Factors Affecting Targeted Sequencing of 353 Nuclear Genes From Herbarium Specimens Spanning the Diversity of Angiosperms

Grace E. Brower¹, James J. Chalmers^{1,2}, Gábor Márton^{1,3}, Alexandros R. Zambidis^{1,4},
Veronica Barlow¹, Silvana Baldo¹, Nicola Baggio¹, Ralf S. Cowan¹, Nara M. J. Dawson¹,
Nerea Delgado¹, Sara L. Edwards¹, Matt C. Edwards¹, Nilsen Edvardsson¹,
Sara Fridolf¹, Ariadne Graft¹, Paul J. Kiersey¹, Lisa Pokorny¹, Ben J. Linder¹, Felix Plass¹
and William J. Baker^{1*}



Herbariomics



REVIEW ARTICLE

INVITED SPECIAL ARTICLE

For the Special Issue: Conducting Botanical Research with Limited Resources: Low-Cost Methods in the Plant Sciences

Strategies for reducing per-sample costs in target capture sequencing for phylogenomics and population genomics in plants

Haley Hale¹, Elliot M. Gardner^{2,4}, Juan Virue⁵, Lisa Pokorny^{6,7}, and Matthew G. Johnson^{1,7}

Manuscript received 8 October 2019; revision accepted 20 December 2019.

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²The Morton Arboretum, Lisle, Illinois 60532, USA.

³Department of Biology, Case Western Reserve University, Cleveland, Ohio 44106, USA.

⁴Singapore Botanic Gardens, National Parks Board, 1 Clump Road, 259569, Singapore.

⁵Royal Botanic Gardens, Kew, Richmond, Surrey TW9 3DS, United Kingdom.

⁶Current address: Centre for Plant Biotechnology and Genomics (CBGP) UPM-INIA, 28223 Pozuelo de Alarcón (Madrid), Spain.

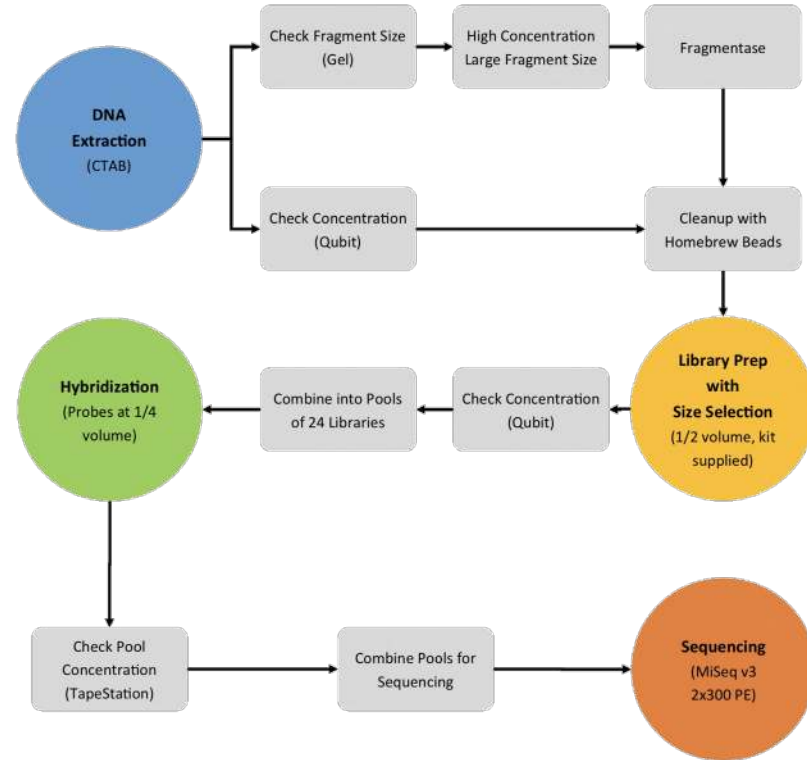
⁷Authors for correspondence: Matt Johnson (mjohnson@ttu.edu), L. Pokorny (lpokorny@ttu.edu).

Citation: Hale H, E M Gardner, J Virue, L Pokorny and M G Johnson. 2020. Strategies for reducing per-sample costs in target capture sequencing for phylogenomics and population genomics in plants. *Applications in Plant Sciences* 8(1): e11337.

doi:10.1002/aps3.11337

The reduced cost of high throughput sequencing and the development of gene sets with wide phylogenetic applicability has led to the rise of sequence capture methods as a plausible platform for both phylogenomics and population genomics in plants. An important consideration in large targeted sequencing projects is the per-sample cost, which can be inflated when using off-the-shelf kits or reagents not purchased in bulk. Here, we discuss methods to reduce per-sample costs in high-throughput targeted sequencing projects. We review the minimal equipment and consumable requirements for targeted sequencing while comparing several alternatives to reduce bulk costs in DNA extraction, library preparation, target enrichment, and sequencing. We consider how each of the workflow alterations may be affected by DNA quality (e.g., fresh vs. herbarium tissue), genome size, and the phylogenetic scale of the project. We provide a cost calculator for researchers considering targeted sequencing to use when designing projects, and identify challenges for future development of low-cost sequencing in non-model plant systems.

KEY WORDS: enzymatic fragmentation; herbariomics; high-throughput workflow implementation; Hyb-Seq; low-cost sequence capture; pooling and multiplexing strategies.



Anatomy of a Plant Radiation



ORIGINAL RESEARCH
published: 20 March 2020
doi: 10.3389/fpls.2020.00258



Reconstructing the Complex Evolutionary History of the Papuanian *Schefflera* Radiation Through Herbariomics

Zhi Qiang Shee^{1,2}, David G. Frodin^{1†}, Rodrigo Cámara-Leret^{1,3,4} and Lisa Pokorny^{1,5,6*}

Assembling the Richest Flora

Article

New Guinea has the world's richest island flora

<https://doi.org/10.1038/s41566-020-2540-5>

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Check for updates

A list of authors and their affiliations appears at the end of the paper.

New Guinea is the world's largest tropical island and has fascinated naturalists for centuries^{1,2}. Home to some of the best preserved ecosystems on the planet³ and to intact ecological gradients—from mangroves to tropical alpine grasslands—that are unmatched in the Asia-Pacific region^{4,5}, it is a globally recognized centre of biological and cultural diversity^{6,7}. So far, however, there has been no attempt to critically catalogue the entire vascular plant diversity of New Guinea. Here we present the first, to our knowledge, expert-verified checklist of the vascular plants of mainland New Guinea and surrounding islands. Our publicly available checklist includes 13,634 species (68% endemic), 1,742 genera and 264 families—suggesting that New Guinea is the most floristically diverse island in the world. Expert knowledge is essential for building checklists in the digital era; reliance on online taxonomic resources alone would have inflated species counts by 22%. Species discovery shows no signs of levelling off, and webinars steps to accelerate botanical research in the ‘Last Unknowns’.

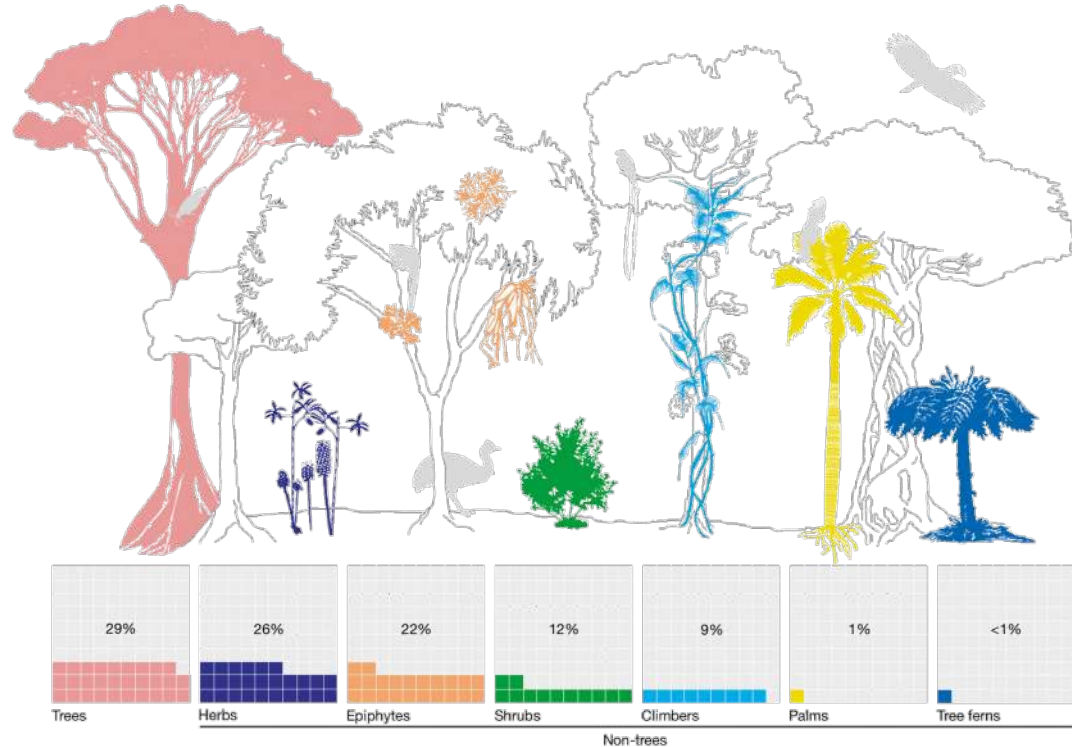
Great uncertainty remains as to the number of New Guinea plant species known to science, with conflicting estimates ranging from 9,000 to 52,000 species^{8–11}. To narrow the range, here we catalogue the entire known vascular flora of New Guinea, including all genera and species of mainland New Guinea and its surrounding islands (hereafter ‘New Guinea’; Fig. 1a, Extended Data Fig. 1). We do so through a large-scale collaborative effort in which 99 plant experts verified the identity of 21,381 taxonomic records in 704,724 specimens (see Methods). Overall, we find that New Guinea supports 13,634 vascular plant species, 1,742 genera and 264 families of vascular plants (Supplementary Tables 1, 2). This suggests that New Guinea is the world's most floristically diverse island, with known vascular plant flora 10% larger than the 12,488 species recorded in Borneo (<http://new-guinea-plant-list.herokuapp.com>, accessed 27 April 2019). New Guinea contains almost three times the 4,598 sporophyte species of Java¹² and 4 times the 9,432 vascular plant species of the Philippines¹³—the only Malayan island regions for which floristic data have been published. The vascular plant flora of New Guinea is divided between two political entities (Fig. 1a). Papua New Guinea, with 10,970 species, has 44 times more species than Indonesian New Guinea (Papua Barat and Papua provinces), which has 2,666. Papua New Guinea also has more species (4,634 versus 3,101) and families (236 versus 248). These differences partly arise from the lower collecting density in Indonesian New Guinea¹⁴ (Fig. 1a). Nevertheless, the order of country rankings in plant diversity is unlikely to change with further collections because Papua New Guinea has a larger area and is far better known than the Indonesian New Guinea¹⁵. Our species total for Papua New Guinea differs markedly from the 29,256 species that are represented in a recent flora of the Global Biodiversity Information Facility¹⁶, and our total number of genera for New Guinea is 28% lower than the 2,457 unvetted genera reported in a previous taxonomic monograph study¹⁷. Together, these differences underscore the need for expert collaboration in the digital era, when we discuss below.

Floristic patterns

Five species-rich families make up 35% of the flora of New Guinea: Orchidaceae (2,856 species), Rubiaceae (974), Eriaceae (438), Poaceae

Endemism

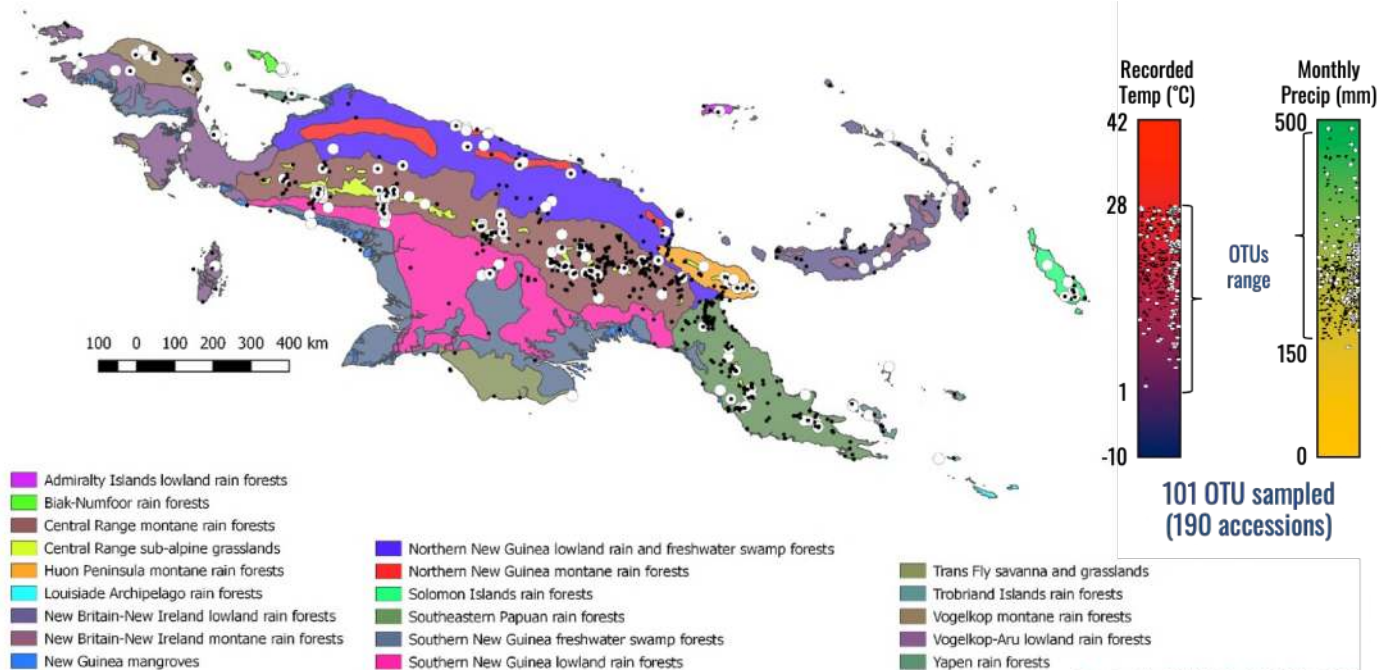
Plant endemism in New Guinea is remarkably high: it is the only Malayan island group with more endemics than more endemic species (9,308 endemic species, 45% of the total). This proportion of endemic species was noted in earlier studies, although these were based on smaller floristic samples^{18,19}. The uniqueness of New Guinea within Malaya may be explained by its greater land surface area and habitat diversity²⁰ in isolation, marking the junction between Malesia, Australia and the Pacific, and its highly complex tectonic history²¹. Geographically, 53% of the endemic species have been found only in Papua New Guinea and 24% occur only in Indonesian New Guinea. Of the total species from Indonesian New Guinea, 4% are endemic, and 38% of the total species from Indonesian New Guinea are endemic. Such high richness of endemic species means that both countries have a unique responsibility for the survival of this irreplaceable biodiversity. Given the trend of plant endemism to increase with elevation²², the conservation of ecosystems along altitudinal gradients is particularly critical. Angiosperms have higher species endemism (76%) than ferns and lycophytes (46%) or gymnosperms (41%). Endemism within families is highly uneven, with just eight angiosperm families comprising 50% of all endemics: Orchidaceae (2,856 endemic species), Rubiaceae (649), Eriaceae (238), Annonaceae (237), Myrsinaceae (255), Gesneriaceae (230), Apocynaceae (196) and Lauraceae (195) (Fig. 1b). The families with the highest proportions of endemism are Eriaceae (98% of species



Umbrella Tree Herbariomics

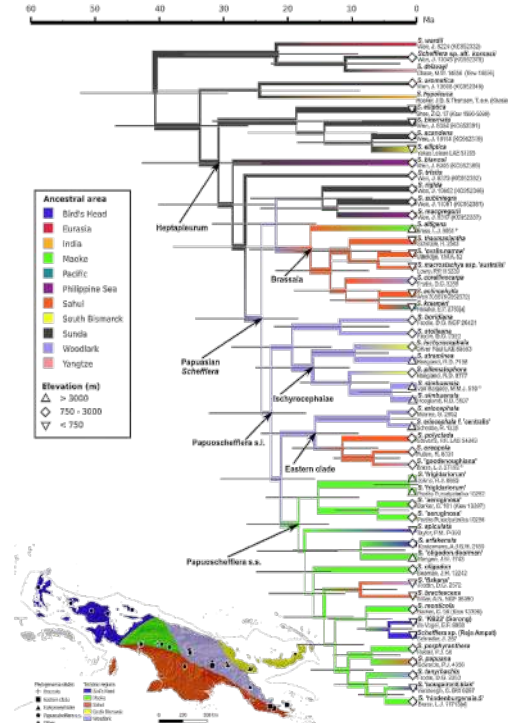
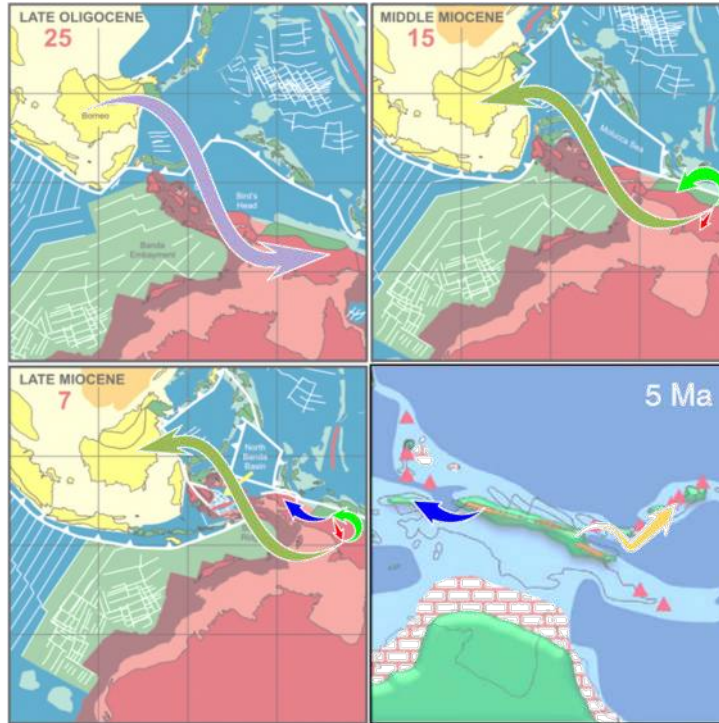


Umbrella Tree Herbariomics



Data from World Wildlife Fund, WorldClim & DG Frodin

Umbrella Tree Geogenomics



A

Phylogenetic tree showing relationships among 100 individuals from 18 populations. The tree is rooted at the top left. Populations are color-coded: Lumak (blue), Pagan (red), and other populations (various colors). A map of Papua New Guinea shows sampling locations. Genetic distances are indicated by numbers along the branches.

B

Bar chart showing genetic differentiation (F_{ST}) across the genome. The y-axis ranges from 0.0 to 1.0. The x-axis shows genomic regions. Two regions are highlighted: Pagan (red) and Lumak (blue).

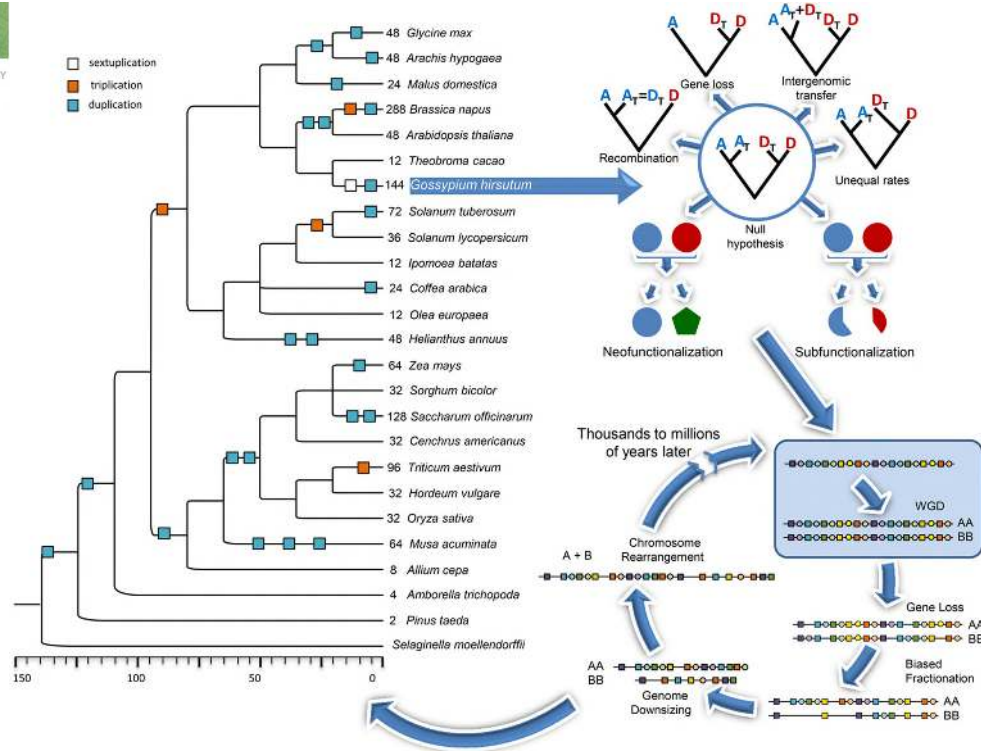
Polyploidy in Plants

NEWS & VIEWS
AMERICAN JOURNAL OF BOTANY

ON THE NATURE OF THINGS: ESSAYS
New Ideas and Directions in Botany

The wondrous cycles of polyploidy in plants¹

Jonathan F. Wendel²



Ploidy Level Estimation

Bioinformatics, 33(16), 2017, 2575–2576
doi: 10.1093/bioinformatics/btx204
Advance Access Publication Date: 5 April 2017
Applications Note

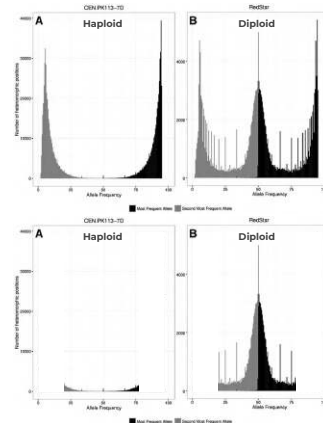
Weiß et al. *BMC Bioinformatics* (2018) 19:122
<https://doi.org/10.1186/s12859-018-2128-z>

BMC Bioinformatics

Genome analysis

ploidyNGS: visually exploring ploidy with Next Generation Sequencing data

Renato Augusto Corrêa dos Santos¹, Gustavo Henrique Goldman² and Diego Mauricio Riaño-Pachón^{1,3,*}



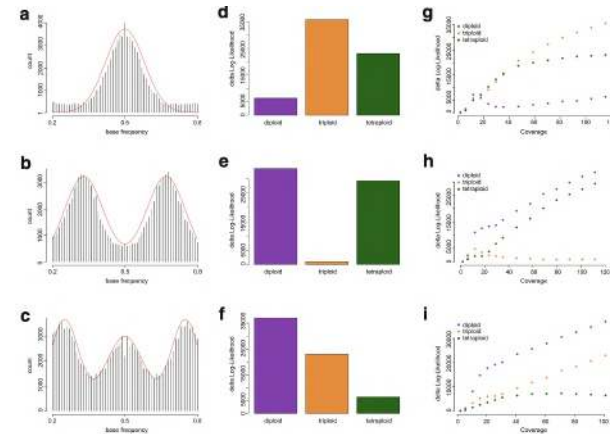
METHODOLOGY ARTICLE

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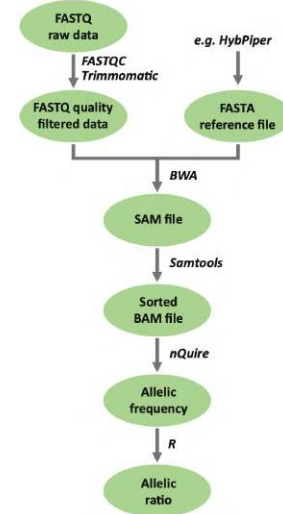


nQuire: a statistical framework for ploidy estimation using next generation sequencing

Clemens L. Weiß¹, Marina Pais², Liliana M. Cano^{2,3}, Sophien Kamoun² and Hernán A. Burbano^{1*}



.....



Dioscoreaceae303

Classification: order Dioscoreales
(Monocot), ~300 species (~4 genera)

Importance: yams (edible tuber),
steroidal saponins (contraceptives), etc.

Available resources:  **1KP**

3 *Dioscorea* & 1 *Tacca* transcriptomes

1 *Oryza sativa* & 1 *Xerophyta viscosa*



MarkerMiner
Locus development made easy

Daicel Arbor Biosciences 

Resulting kit:

260 ~single copy nuclear orthologs

43 genes of agronomic interest

~442 Kbp matrix

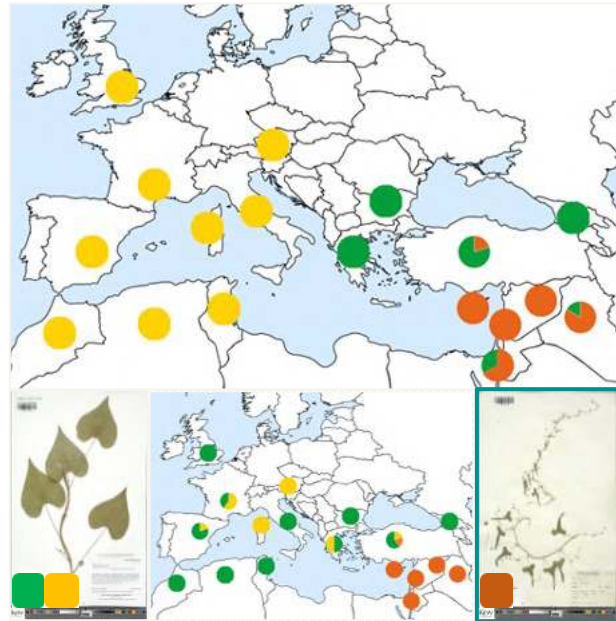
~17K 120-mer 3x tiled probes



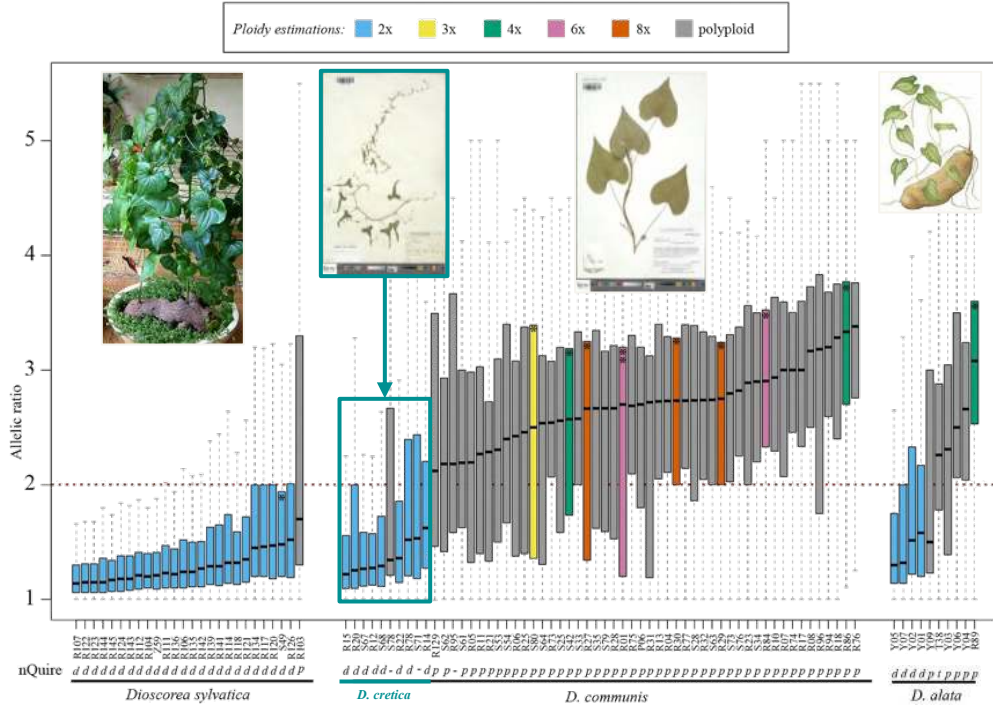
**A customized nuclear target enrichment approach for
developing a phylogenomic baseline for *Dioscorea* yams
(Dioscoreaceae)**

Marybel Soto Gomez^{1,2*} , Lisa Pokorny³ , Michael B. Kantar⁴ , Félix Forest⁵ , Ilija J. Leitch⁶ , Barbara Gravendeel^{MA7} ,
Paul Wilkin¹ , Sean W. Graham^{1,4} , and Juan Viruel⁶ 

Ploidy Level in Yams



D. cretica



Ploidy Level in Yams

Annals of Botany 131: 635–654, 2023
https://doi.org/10.1093/aob/mcad018, available online at wwwacademic.oup.com/aob

ANNALS OF
BOTANY

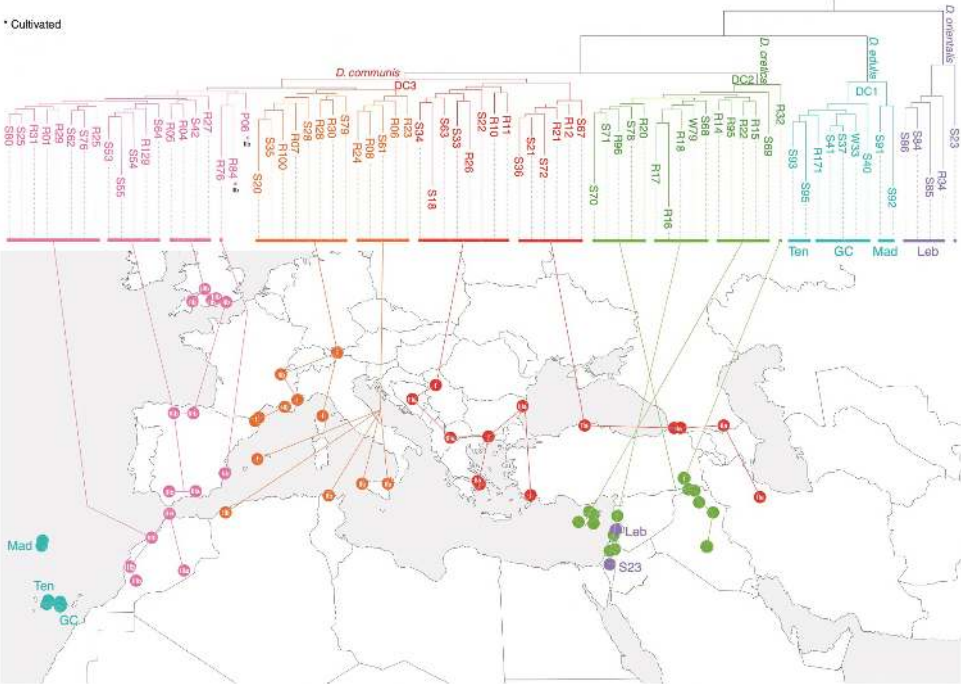
Genomic, spatial and morphometric data for discrimination of four species in the Mediterranean Tamus clade of yams (*Dioscorea*, Dioscoreaceae)

Miguel Campos^{1,2,3}, Emma Kelley¹, Barbara Gravendeel^{1,4}, Frédéric Médail⁵, J. M. Maarten Christenhusz¹, Michael F. Fay^{1,2}, Pilar Catalán^{1,4}, Ila J. Leitch¹, Félix Forest¹, Paul Wilkin¹ and Juan Viruel^{1,6}*

¹Royal Botanic Gardens, Kew, Richmond TW9 3DS, UK, ²Department of Plant Biology and Ecology, University of Seville, 41012, Spain, ³Universidad de Zaragoza-Escuela Politécnica Superior de Huesca, 22071, Huesca, Spain, ⁴Naturalis Biodiversity Center, Leiden 2333 CR, The Netherlands, ⁵Radboud Institute for Biological and Environmental Sciences, RIBES 6500 GL, Nijmegen, The Netherlands, ⁶Institut Méditerranéen de Biodiversité et d'Écologie marine et continentale (IMBE), Aix Marseille University, Avignon University, CNRS, IRD, Campus Aix, Technopôle de l'Environnement Arbois-Méditerranée, F-13545 Aix-en-Provence cedex 4, France, ⁷School of Biological Sciences, University of Western Australia, Crawley, WA 6009, Australia and ⁸Grupo de Bioquímica, Biofísica y Biología Computacional (BIFI, UNIZAR), Unidad Asociada al CSIC, Zaragoza 50018, Spain

*For correspondence. E-mail j.viruel@kew.org, juanviruel@gmail.com

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Electronically published: 22 January 2023



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Cortinariaceae193

Classification: order Agaricales
(Basidiomycota), ~3K species (1 genus)

Importance: ectomycorrhizal fungi

Available resources:

4 *Cortinarius* assemblies (ABYSS)
208 Agaricales single-copy orthologs
genes (SCOGs) & exonerate
Daicel Arbor Biosciences 

Resulting kit:

188 nuclear SCOGs
RPB1, RPB2, MCM7, GPD, & TEF1
~67.5 Kbp matrix
~20K 120-mer 2x tiled probes

Fungal Diversity
<https://doi.org/10.1007/s13225-022-00499-9>



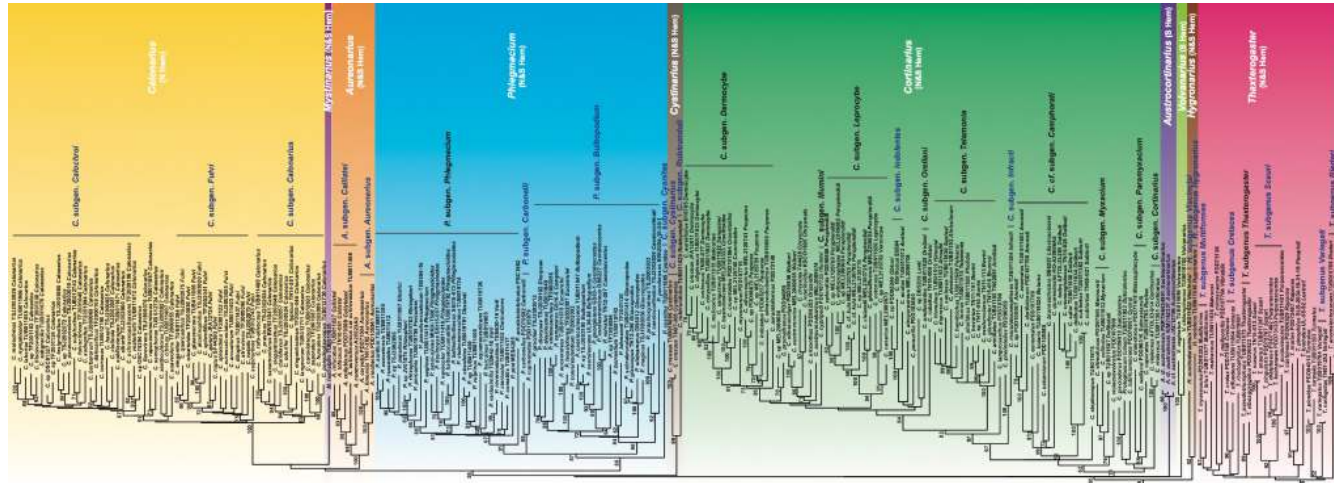
Taming the beast: a revised classification of *Cortinariaceae* based on genomic data

Kare Liimatainen¹ · Jan T. Kim² · Lisa Pokorny^{1,3} · Paul M. Kirk⁴ · Bryn Dentinger⁵ · Tuula Niskanen^{1,6}

Received: 20 October 2021 / Accepted: 20 January 2022
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Table 3 Species diversity, distribution, and summary of the selected morphological characteristics of the 10 genera of *Cortinariaceae*

Genus	Generic syno- nyms	Estimated no. of spe- cies	Subgen./sec- tion no	Distribution	Cystidia	Stipitocarpic (S)/Pileo- carpic (P)	Pileipellis simplex/ duplex	Sequestrate sp	Appearance of the agaricoid basidiomata				Rozitoid/ cuphocy- boid
									Myxacioid	Phlegmac	Telamonioid	Cortinarioid	
<i>Cortinarius</i>	11	>2000	11/130*	N+S	(X)	S/(P)	Duplex (+ simpl.)	X	X	(X)	X	X	X
<i>Phlegmacium</i>	3	>200	4/23	N (+S)		P/S	Duplex (+ simpl.)	X		X	(X)		
<i>Thaxtero- gaster</i>	3	>200	6/22	(N+) S		S/P	Duplex	X	X	X			
<i>Calonarius</i>	—	~200	3/14	N		P	Simplex	X		X			
<i>Aureonarius</i>	—	~25	2/3	(N+) S		S	Duplex					X	
<i>Cystinarius</i>	—	~10	2/2	N+S	X	S	Duplex			X		(X)	
<i>Volvanarius</i>	—	~10	1/1	S	(X)	P	Duplex	X		X			
<i>Hygromarius</i>	—	~10	2/2	(N+) S		S	Duplex				X		
<i>Austrocorti- narius</i>	—	<5	1/1	S		S	Duplex			X			
<i>Mystinarius</i>	—	<5	1/1	N+S		S	Duplex		X	X			



Cantharellales526

Classification: Agaricomycetes
(Basidiomycota), ~630 spp. (~40 gen.)

Importance: ectomycorrhizal & edible

Available resources:

10 DOE-JGI genomes (1KFG)

PHYling & UFGG SCOGs

get_homologues, HybPiper, AMAS

Daicel Arbor Biosciences 

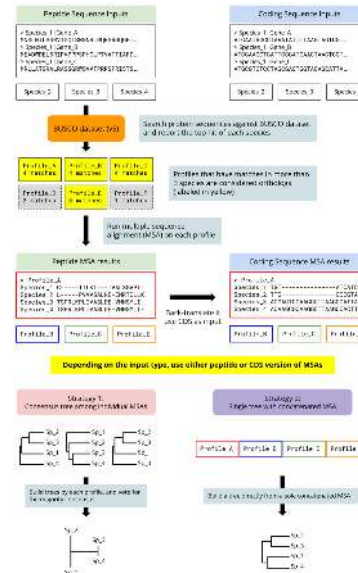
Resulting kit:

523 nuclear SCOGs

RPB1&2, EF, & MCM7

~7.5 Mbp matrix

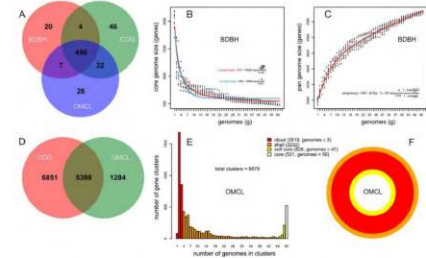
~55K 120-mer 1x tiled probes



GET_HOMOLOGUES: a versatile software package for pan-genome analysis

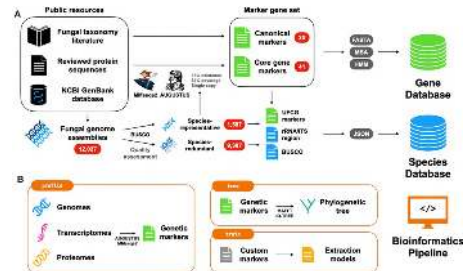
Code [Genomes](#) [Bioinformatics](#) [GitHub](#) [DOI](#)

This software is maintained by Bruno Contreras-Moreira (bcontreras at eead.csic.es) and Pablo Vinuesa (pvinuesa at ccg.unam.mx). The original version, suitable for bacterial genomes, was described in:



Contreras-Moreira B, Vinuesa P (2013) Appl. Environ. Microbiol. 79:7696-7701

Vinuesa P, Contreras-Moreira B (2015) Methods in Molecular Biology Volume 1231, 203-232



Cantharellales526

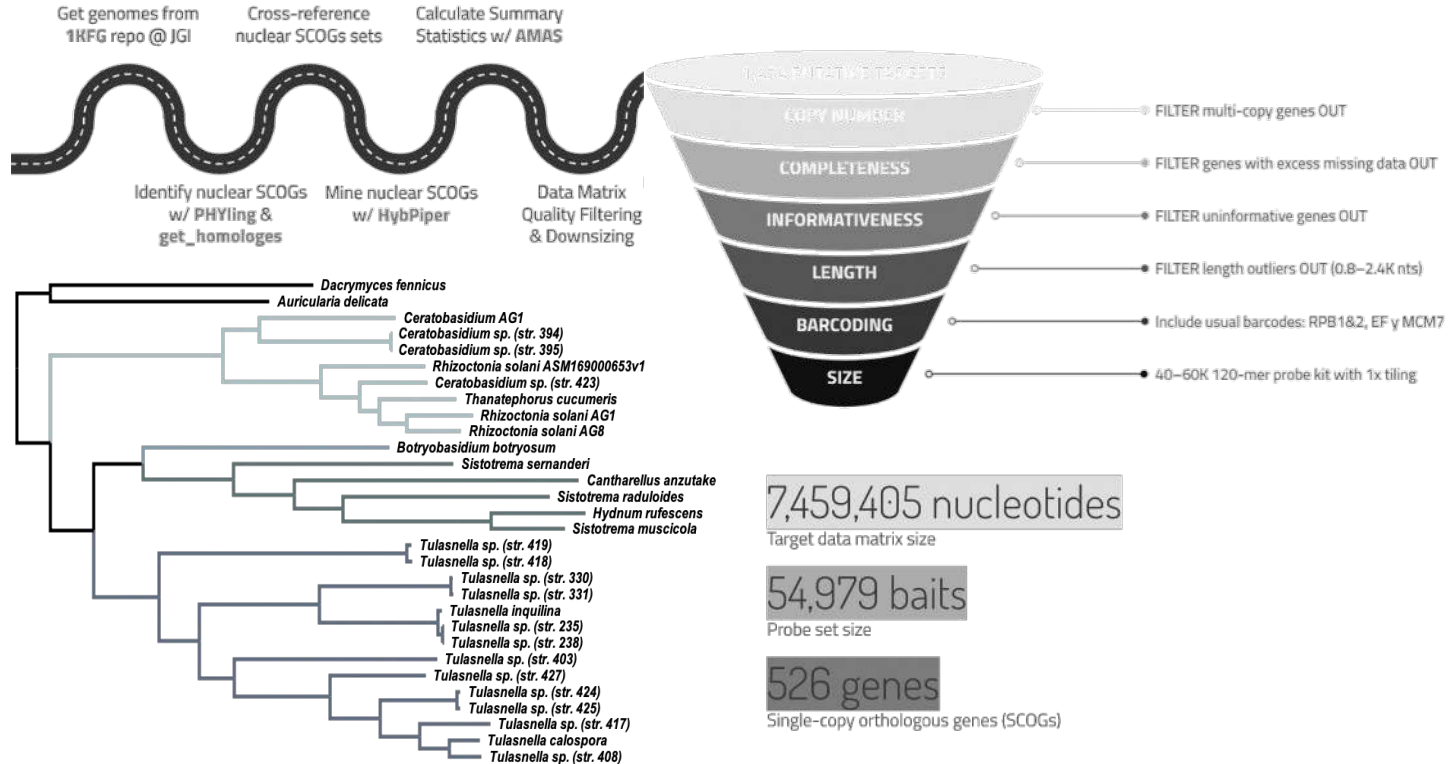


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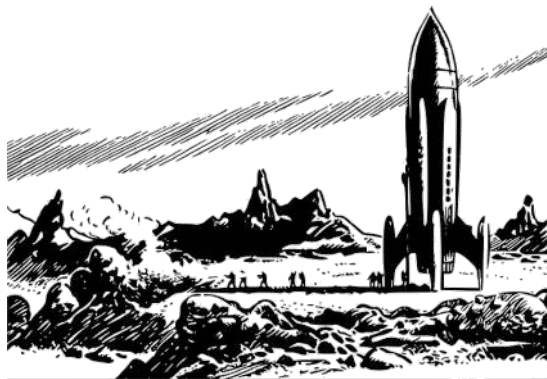
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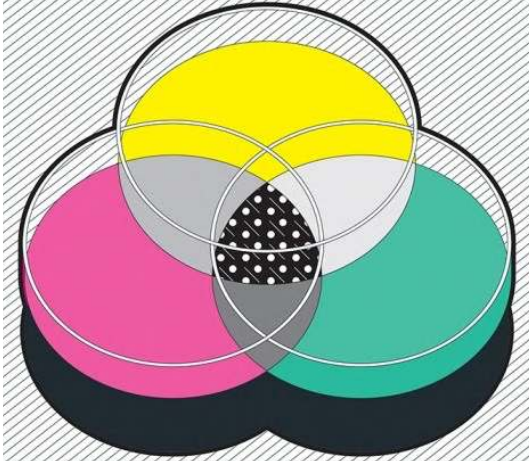
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Exploring the Known Unknowns and the
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Matter)



Mesoevolutionary Approaches

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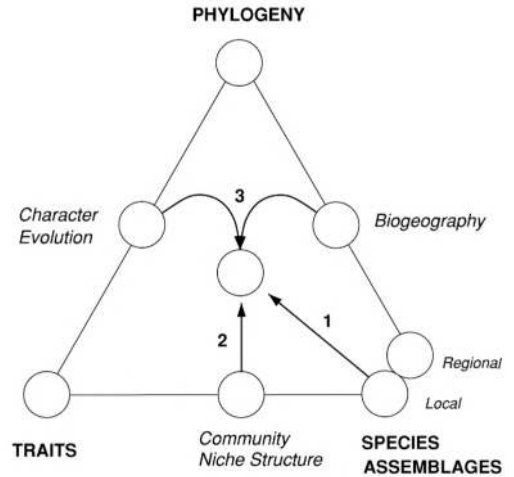
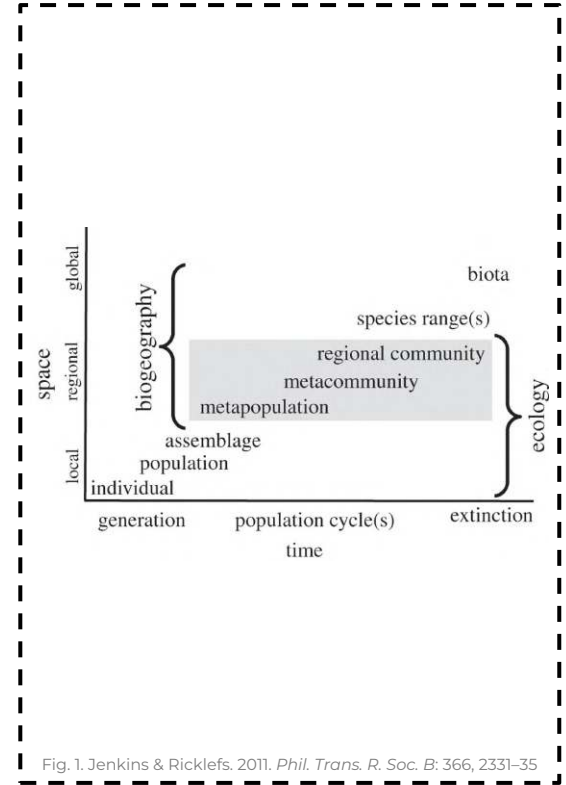
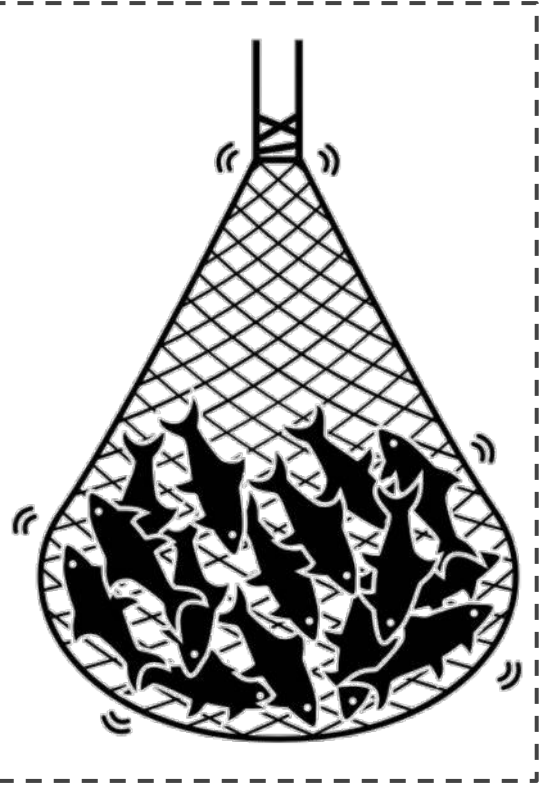


Fig. 1. Webb et al. 2002. *Annu. Rev. Ecol. Syst.* 33:475–505.

Community Phylogenomics

At the Intersection of Biogeography and Ecology





Thanks!

Questions?

pokorny@rjb.csic.es
@Calypstrochaeta

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