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Data Article

Genome assembly and phylogenomic data analyses using plastid data: Contrasting species tree estimation methods



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ABSTRACT

Phylogenomics has become increasingly popular in recent years mostly due to the increased affordability of next generation sequencing techniques. Phylogenomics has sparked interest in multiple fields of research, including systematics, ecology, epidemiology, and even personalized medicine, agriculture and pharmacy. Despite this trend, it is usually difficult to learn and understand how the analyses were done, how the results were obtained, and most importantly, how to replicate the study. Here we present the data and all of the code utilized to perform phylogenomic inferences using plastome data: from raw data to extensive phylogenetic inference and accuracy assessment. The data presented here utilizes plastome sequences available on GenBank (accession numbers of 94 species are available below) and the code is also available at https://github.com/deisejpg/rosids. Gonçalves et al. is the research article associated with the data analyses presented here.

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Specifications table

Subject area	Biology				
More specific subject	lore specific subject Systematics of angiosperms, plastome evolution				
area					
Type of data	Table, PHYLIP, NEXUS, and MARKDOWN formatted files				
How data was acquired	Illumina HiSeq. 2500 and Illumina HiSeq 4000				
Data format	Raw and Analyzed				
Experimental factors	Total genomic DNA isolated and sequenced from fresh or silica dried leaf tissue of sampled species of rosids and genome sequences from NCBI				
Experimental features	Illumina reads preprocessing, genome assembly, phylogenetic inference, topology tests, and phylogenetic signal assessment				
Data accessibility	Within this article and at https://github.com/deisejpg/rosids				
Related research	Deise J.P. Gonçalves, Beryl B. Simpson, Edgardo M. Ortiz, Gustavo H. Shimizu, Robert K. Jansen				
article	Incongruence between gene trees and species trees and phylogenetic signal variation in plastid genes				
	Molecular Phylogenetics and Evolution				
	10.1016/j.ympev.2019.05.022				

Value of the data

The present data provides details about phylogenomic analysis using a set of well-documented pipelines covering
analyses from preprocessing Illumina reads to inferring and testing phylogenies using multiple methods of
phylogenetic inference

• The data introduce the practical use of multispecies coalescent methods using plastid protein-coding genes and could be adjusted and used with molecular data from different molecular markers and organisms

 Accessibility to scripts utilized and data files containing the alignments and trees will enhance the replication of the analyses presented

1. Data

A dataset comprising 78 plastid protein-coding genes of 94 species of rosids is presented in Table 1. Here we present all the code used in the analysis of this dataset [1], including the scripts used to quality filter, assemble, extract regions of interest, and perform phylogenomic analysis, in a series of tutoriallike files: I. Genome assembly; II. Phylogenetic Analysis; III. Tree space; IV. Phylogenetic Signal. Part of the data was obtained from GenBank (http://www.ncbi.nlm.nih.gov/genbank). Data for and 27 species from groups of rosids that lacked the information on the database were generated using Illumina HiSeq. A total of 657,471,631 million paired-end reads with an average length of 150 bp was generated (Table 2). Despite the interest on extracting and using only the genes from the plastome, the pipeline for genome assembly presented here also separates contigs from the three cellular genomic compartments with a potential for use used in studies that target not only the plastome, but also mitochondria, nuclear ribosomal DNA, and other nuclear markers. The next set of tutorial-like markdown files present the code utilized for preparing alignments and for inferring phylogenies using an array of strategies of data partition and methods of phylogenetic inference. The code used to explore the similarities/dissimilarities between topologies and for the phylogenetic signal calculation is also presented.

2. Experimental design, materials and methods

2.1. Data preprocessing and genome assembly

For the 27 samples for which data were generated, leaf tissue was ground and total genomic DNA was isolated using DNeasy Plant Mini Kit (Qiagen) according to the manufacturer's protocol or a modified version of [2] described in Ref. [3]. The DNA was quantified using a Qubit Fluorometric Quantitation (Thermo Fisher) instrument and was sequenced at the Genome Sequencing and Analysis

Table 1

Classification according to APG IV (2016) of samples used in the study (89 samples of rosids, considered here as fabids + malvids, and five of outgroup), voucher information of newly sequenced plastomes and GenBank accession numbers. Bold font indicates plastid genome sequences generated in this study.

Order	Family	Species	Voucher ID	GenBank Accession Numbers
Ingroup				
Brassicales	Brassicaceae	Brassica napus		NC 016734
Brassicales	Caricaceae	Carica papava		NC 010323
Brassicales	Moringaceae	Moringa oleifera	CONN-129179	MK726020
Brassicales	Salvadoraceae	Azima tetracantha	CONN00225893	MK726028
Celastrales	Celastraceae	Euonymus japonicus		NC_028067.1
Cucurbitales	Cucurbitaceae	Citrullus lanatus		NC_032008
Cucurbitales	Cucurbitaceae	Cucumis hystrix		NC_023544
Cucurbitales	Cucurbitaceae	Gynostemma pentaphyllum		NC_029484
Fabales	Fabaceae	Cicer arietinum		NC_011163
Fabales	Fabaceae	Inga leiocalycina		NC_028732
Fabales	Fabaceae	Lupinus luteus		NC_023090
Fabales	Polygalaceae	Polygala alba	TEX-DJPG731	MK726019
Fagales	Betulaceae	Ostrya rehderiana		NC_028349
Fagales	Fagaceae	Castanea mollissima		NC_014674
Fagales	Juglandaceae	Juglans regia		NC_028617
Geraniales	Francoaceae	Francoa sonchifolia		NC_021101
Geraniales	Francoaceae	Melianthus villosus		NC_023256
Geraniales	Francoaceae	Viviania marifolia		NC_023259
Geraniales	Geraniaceae	Erodium rupestre		NC_030719
Geraniales	Geraniaceae	Geranium palmatum		NC_014573
Geraniales	Geraniaceae	Hypseocharis bilobata		NC_023260
Geraniales	Geraniaceae	Monsonia speciosa		NC_014582
Geraniales	Geraniaceae	Pelargonium alternans		NC_023261
Malpighiales	Chrysobalanaceae	Chrysobalanus icaco		NC_024061
Malpighiales	Chrysobalanaceae	Hirtella racemosa		NC_024060
Malpighiales	Erythroxylaceae	Erythroxylum novogranatense		NC_030601
Malpigniales	Eupnorbiaceae	Ricinus communis	TEV DIDC002	NC_016736
Malpighiales	Saligagoago	Gaiphimia angustijona	TEX-DJPG803	MK726010
Malvalos	Malvacoao	Cossumium turnari		NC_026825
Malvalos	Malvaceae	Hibicaus springus		NC_026000
Malvales	Malvaceae	Theobroma cacao		NC_020505
Malvales	Thymelaeaceae	Aquilaria sinensis		NC 029243
Murtales	Alzateaceae	Alzatea verticillata	K-TNV548	MK726006
Myrtales	Combretaceae	Laguncularia racemosa	CONN00225898	MK726017
Myrtales	Combretaceae	Terminalia guvanensis	UEC-GHS1070	MK726027
Myrtales	Lythraceae	Heimia anetala	CONN00225896	MK726012
Myrtales	Lythraceae	Lagerstroemia guilinensis		NC 029885
Myrtales	Lythraceae	Lagerstroemia fauriei		NC 029808
Myrtales	Lythraceae	Lagerstroemia indica		NC 030484
Myrtales	Melastomataceae	Blakea schlimii		NC_031877
Myrtales	Melastomataceae	Henriettea barkeri		NC_031880
Myrtales	Melastomataceae	Memecylon pauciflorum	K-TNV679	MK726029
Myrtales	Melastomataceae	Miconia dodecandra		NC_031882
Myrtales	Melastomataceae	Rhexia virginica		NC_031886
Myrtales	Melastomataceae	Tibouchina urvilleana	CONN00225897	MK726030
Myrtales	Myrtaceae	Allosyncarpia ternata		NC_022413
Myrtales	Myrtaceae	Corymbia eximia		NC_022409
Myrtales	Myrtaceae	Eucalyptus globulus		NC_008115
Myrtales	Myrtaceae	Eugenia uniflora		NC_027744
Myrtales	Myrtaceae	Heteropyxis natalensis	K-MFF s.n.	MK726014
Myrtales	Myrtaceae	Psidium guajava		NC_033355
Myrtales	Myrtaceae	Stockwellia quadrifida		NC_022414
Myrtales	Myrtaceae	Xanthostemon chrysanthus	K-TNV684	MK726024
Myrtales	Onagraceae	Ludwigia octovalvis		NC_031385
Myrtales	Unagraceae	Uenothera argillicola		NC_010358

(continued on next page)

Table 1 (continued)

Order	Family	Species	Voucher ID	GenBank Accession
				Numbers
Myrtales	Onagraceae	Oenothera grandiflora		NC_029211
Myrtales	Onagraceae	Oenothera oakesiana		NC_029212
Myrtales	Onagraceae	Oenothera villaricae		NC_030532
Myrtales	Penaeaceae	Saltera sarcocolla	CONN00225892	MK726025
Myrtales	Vochysiaceae	Callisthene erythroclada	UEC-DG439	MK726008
Myrtales	Vochysiaceae	Erisma bracteosum	UEC-GHS937	MK726009
Myrtales	Vochysiaceae	Erismadelphus exsul	BRLU-SB751	MK726007
Myrtales	Vochysiaceae	Korupodendron	BRLU-SB3491	MK726013
-	-	songweanum		
Myrtales	Vochysiaceae	Qualea grandiflora	UEC-DG382	MK726022
Myrtales	Vochysiaceae	Ruizterania albiflora	UEC-TM258	MK726023
Myrtales	Vochysiaceae	Salvertia convallariodora	TEX-DJPG569	MK726026
Myrtales	Vochysiaceae	Vochysia acuminata	TEX-DJPG150	MK726031
Oxalidales	Oxalidaceae	Averrhoa carambola		NC_033350
Oxalidales	Oxalidaceae	Oxalis drummondii	TEX-DJPG722	MK726021
Rosales	Cannabaceae	Cannabis sativa	-	NC_026562
Rosales	Elaeagnaceae	Elaeagnus macrophylla		NC_028066
Rosales	Moraceae	Ficus racemosa		NC_028185
Rosales	Moraceae	Morus indica		NC_008359
Sapindales	Sapindaceae	Acer morrisonense		NC_029371
Sapindales	Anacardiaceae	Anacardium occidentale		NC_035235.1
Sapindales	Anacardiaceae	Mangifera indica		NC_035239.1
Sapindales	Burseraceae	Boswellia sacra		NC_029420
Sapindales	Meliaceae	Azadirachta indica		NC_023792
Sapindales	Rutaceae	Citrus aurantiifolia		NC_024929
Sapindales	Rutaceae	Zanthoxylum piperitum		NC_027939
Sapindales	Sapindaceae	Dimocarpus longan	UCONN-201400014	MK726005
Sapindales	Sapindaceae	Dipteronia sinensis		NC_029338
Sapindales	Sapindaceae	Litchi chinensis		NC_035238.1
Sapindales	Sapindaceae	Sapindus mukorossi		NC_025554
Vitales	Vitaceae	Tetrastigma hemsleyanum		NC_029339
Vitales	Vitaceae	Vitis vinifera		NC_007957
Zygophyllales	Krameriaceae	Krameria bicolor	TEX-JN14-09-03-1	MK726015
Zygophyllales	Krameriaceae	Krameria lanceolata	TEX-JN6-IV-2015-1	MK726016
Zygophyllales	Zygophyllaceae	Guaiacum angustifolium	TEX-BBS20-IV-2015-1	MK726011
Zygophyllales	Zygophyllaceae	Larrea tridentata	TEX-JN14-08-24-1	MK726018
Outgroup				
Caryophyllales	Caryophyllaceae	Silene capitata		NC_035226
Apiales	Araliaceae	Aralia undulata		NC_022810
Solanales	Solanaceae	Nicotiana tabacum		NC_001879.2

Facility (GSAF) at The University of Texas at Austin. Two species were kindly provided by The Royal Botanic Gardens, Kew, DNA bank (https://www.kew.org/data/dnaBank/).

Once the reads were available, the genome assembly pipeline was used to remove adaptors and PHIX, for quality trimming, and for genome assembly.

2.2. Phylogenetic inference

After gathering sequences of plastid protein-coding genes, the alignments and phylogenetic inference were performed. The code used to prepare the alignments using MAFFT [4] and MACSE [5] as well as the scripts used to infer phylogenies using Maximum Likelihood (ML), IQ-TREE [6], and Multispecies Coalescent (MSC) methods, SVDquartets [7], and ASTRAL-II [8] is presented in phylogenetic analysis pipeline.

2.3. Calculating distances of tree topologies and phylogenetic signal

Commented scripts present how the inferred phylogenies were further explored. First, Robinson-Foulds and Kendall-Colijn algorithms implemented in the R package TREESPACE [9] were used to

Table 2

Summary of output for Illumina sequencing of the 23 complete and the 4 draft plastomes.

Species	Total # Reads	Plastid reads	Average fold coverage ^a
Laguncularia racemosa Terminalia guvanensis	26,332,422	992,869	758.64
Heimia anetala	27,057,568	3/80 002	3630.02
Memocylon nauciflorum	27,037,300	1 776 470	2078 67
Tibouchina urvilleana	51 945 408	1,770,470	1321 25
Heteropyyis natalensis	23 511 880	437 572	620 57
Saltera sarcocolla	42 509 058	1 975 497	1589.48
Callisthene erythroclada	13 523 069	1,375,457	1679.93
Frisma bracteosum	28 714 784	740 005	872.25
Qualea grandiflora	14 026 674	315 666	361 56
Korupodendron songweanum	26 380 860	621 934	697 51
Ruizterania albiflora	13,416,345	346.923	395.21
Salvertia convallariodora	13.719.886	1.131.571	1262.57
Vochysia acuminata	14,190,315	1,404,610	1588.55
Azima tetracantha	19,768,664	588,345	676.25
Moringa oleifera	35,924,836	3,914,845	3054.87
Dimorcarpus longan	39,914,336	1,125,881	870.47
Galphimia angustifolia	20,773,804	1,235,549	1449.65
Oxalis drummondii	30,898,958	1,231,010	1698.62
Krameria bicolor	20,882,876	1,032,768	1236.57
Krameria lanceolata	20,840,984	191,073	206.57
Guaiacum angustifolium	27,224,600	131,532	143.81
Larrea tridentata	23,104,094	2,094,664	2797.51
Alzatea verticillata (reads mapped to S. sarcocolla)	11,665,088	51,656	61.91
Xanthostemon chrysanthus	9,809,306	51,304	58.28
(reads mapped to <i>H. natalensis</i>)			
Polygala alba (reads mapped to G. angustifolia)	30,134,340	125,019	101.66
Erismadelphus exsul (reads mapped to K. songweanum)	13,270,682	286,659	368.03

^a Coverage calculation was performed in bbmap (BBTools package, available at https://igi.doe.gov/data-and-tools/bbtools/) with the option "covstats". Reference index was built with the 23 complete plastomes with each representing a scaffold. The average fold coverage was calculated by mapping reads to the reference index and values were taken from the scaffold correspondent to each species. For species with incomplete plastomes (marked in bold) we used the closely related species as the scaffold indicated within the parentheses.

visualize the distances of species trees and between species trees and gene trees inferred. The code used is available at tree space. Lastly, five taxa from different taxonomic levels that had alternative placements were selected for a set of measurements of gene-wise and site-wise log-likelihood support of alternative topologies phylogenetic signal.

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Conflict of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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