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Global Brassicaceae phylogeny based on filtering of 1,000-gene dataset

Highlights

- An unparalleled Brassicaceae phylogeny covering nearly all 349 genera is presented
- Cytonuclear discordance is omnipresent and a likely sign of rampant hybridization
- The family originated during the late Eocene to late Oligocene
- Results are used to come up with a new family classification

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In brief

The Brassicaceae family contains the model plant *Arabidopsis thaliana* and many important crop species. Surprisingly, relationships within the family were poorly known. Hendriks et al. present an unprecedented family phylogeny with representatives from nearly all 349 genera. These results will boost fundamental and applied plant biological research.



Article

Global Brassicaceae phylogeny based on filtering of 1,000-gene dataset

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SUMMARY

The mustard family (Brassicaceae) is a scientifically and economically important family, containing the model plant *Arabidopsis thaliana* and numerous crop species that feed billions worldwide. Despite its relevance, most phylogenetic trees of the family are incompletely sampled and often contain poorly supported branches. Here, we present the most complete Brassicaceae genus-level family phylogenies to date (Brassicaceae Tree of Life or BrassiToL) based on nuclear (1,081 genes, 319 of the 349 genera; 57 of the 58 tribes) and plastome (60 genes, 265 genera; all tribes) data. We found cytonuclear discordance between the two, which is likely a result of rampant hybridization among closely and more distantly related lineages. To evaluate the impact of such hybridization on the nuclear phylogeny reconstruction, we performed five different gene sampling routines, which increasingly removed putatively paralog genes. Our cleaned subset of 297 genes revealed high support for the tribes, whereas support for the main lineages (supertribes) was moderate. Calibration based on the 20 most clock-like nuclear genes suggests a late Eocene to late Oligocene origin of the family. Finally, our results strongly support a recently published new family classification, dividing the family into two subfamilies (one with five supertribes), together representing 58 tribes. This includes five recently described or re-established tribes, including Arabidopsidae, a monogeneric tribe accommodating *Arabidopsis* without any close relatives. With a worldwide community of thousands of researchers working on Brassicaceae and its diverse members, our new genus-level family phylogeny will be an indispensable tool for studies on biodiversity and plant biology.

INTRODUCTION

The mustard family (Brassicaceae) is a globally distributed, medium-sized plant family (~4,000 species) with huge economic

and scientific impact, characterized by high morphological diversity^{1–5} (Figure 1). The family contains numerous species grown for food and biofuel (cabbage, rapeseed) as well as many model species (*Arabidopsis thaliana*, *Arabis alpina*,



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Brassica spp., and *Capsella* spp.).^{1,6} Moreover, the overwhelming availability of genetic tools and resources in these species make Brassicaceae an ideal model family in flowering plants. These tools have facilitated, among other things, plant developmental studies that (1) disentangle genotype-phenotype interactions,^{7,8} (2) investigate impacts of whole-genome duplications (WGDs) at different timescales,^{9,10} and (3) unravel the evolutionary variation in metabolic pathways, leading to a huge diversity in natural products.¹¹ In many of these comparative studies that go beyond model species, a robust evolutionary framework is required, ideally encompassing the ~4,000 currently accepted species of Brassicaceae that are divided among 349 genera and 50–60 tribes.¹² This is also true for studies on crop wild relatives aiming to introgress desirable traits (e.g., drought tolerance and disease resistance) into crops using plant breeding.^{13–15} Therefore, the evolution of Brassicaceae has been the subject of study for a long time.^{2,3,16–23} However, a robust, densely sampled Brassicaceae Tree of Life (hereafter named BrassiToL) remains lacking.

Early phylogenetic inferences relying on flower and fruit morphology^{25–27} were often misled by rampant convergent evolution.^{28–30} Subsequently, the use of few molecular markers generated poorly supported phylogenies,^{17,31} traditionally attributed to an early rapid radiation²⁰ and multiple WGD and hybridization events.¹ More recent family-wide phylogenetic studies based on multiple molecular markers offered support for three,³² four,¹ or five^{3,22,33–35} main lineages in addition to tribe Aethionemeae, which is sister to all other Brassicaceae. However, several evolutionary events in the family keep challenging the reconstruction of the BrassiToL backbone,^{10,36–40} including incomplete lineage sorting and introgression,⁴¹ and (ancient)

inter-tribal and inter-lineage hybridization, commonly followed by post-polyploid diploidization.^{42,43}

Here, we present results from the largest global Brassicaceae phylogenetic study to date including nearly all genera, with the following objectives: (1) reconstruct the most complete and robust family-wide BrassiToL based on nuclear genomic data and plastome data, (2) investigate the influence of paralogous genes and polyploid species on phylogeny reconstruction, (3) re-evaluate the temporal origin of the Brassicaceae, its main lineages (subfamilies and supertribes), and tribes, and (4) provide an updated taxonomic delimitation of these lineages and tribes. Our improved phylogenetic framework will further develop Brassicaceae as the prime model family in flowering plants.

RESULTS

Unparalleled family-wide sampling

Our ingroup sampling for the nuclear analyses included 380 samples, covering 375 species, 319 of the 349 currently accepted genera (91%) and 57 of the 58 accepted tribes following the latest taxonomic revision of German et al.²⁴ (excluding Hilliellae; [Data S1A](#) and [S1B](#)). We added 23 outgroup species representing all Brassicales families and 14 former Brassicaceae genera now synonymized with accepted genera. Three additional genera (*Hilliella*, *Onuris*, and *Thelypodium*) were included in the plastome dataset ([Data S1C](#)).

We recovered 1,081 nuclear genes ([Data S1D](#); [Methods S1A](#)) across 397 samples ([Data S1E](#)) from target capture sequencing of 764 Brassicaceae-specific genes³ (hereafter B764) and 353 Angiosperm-wide genes^{44,45} (hereafter A353), following a “mixed

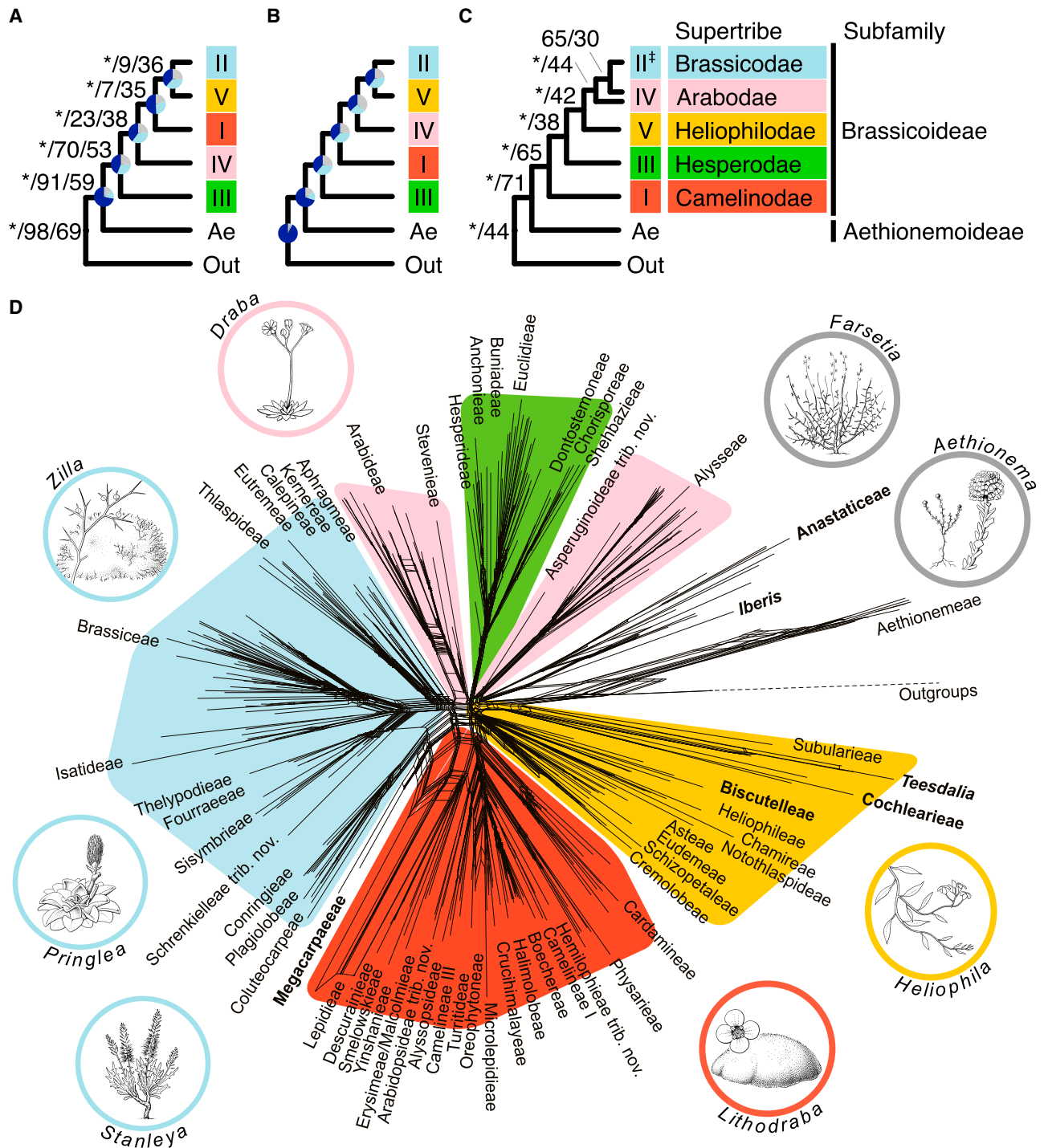


Figure 1. Overview of the main mustard family relationships from different phylogenetic reconstruction approaches and gene sampling routines

Color group is the main core Brassicaceae lineages (I–V) described by Nikolov et al.,³ with recently proposed names for subfamilies and supertribes following German et al.²⁴ See also Figure S3.

(A) Cladogram of main lineages as recovered from nuclear stricter routines (strict, superstrict, superstrict by tribe, and superstrict excluding hybrids) using either supermatrix ML or coalescent-based approaches (ASTRAL-III), with node support from the supermatrix ML approach on the superstrict excluding hybrids routine (BS/gCF/sCF; * indicates 100%) and the ASTRAL-III approach on the superstrict routine (pie charts represent quartet scores: dark blue for the first quartet, light blue for the first alternative, and gray for the second alternative).

(B) Cladogram of main lineages as recovered from nuclear inclusive routine using coalescent-based approach (using both ASTRAL-III and ASTRAL-Pro) with node support from the ASTRAL-III approach on the inclusive routine (colors as in previous panel).

(legend continued on next page)

baits” methodology.⁴⁶ For B764, we recovered 91% of the total target length of 888,392 bp (mean across all samples), and 84% of the 728 unique genes targeted (36 genes in common with the A353 kit were first removed). For A353, mean values were 83% of the total target length of 294,516 bp and 80% of 353 genes targeted. Interestingly, gene recovery hardly decreased with sample age, and we successfully obtained hundreds of nuclear genes even from pre-1900 samples (Methods S1B).

Unparalleled global nuclear Brassicaceae phylogeny

We used HybPhaser v2.0⁴⁷ to define five “sampling routines” (“inclusive”: 1,018 genes, “strict”: 1,013 genes, “superstrict”: 297 genes, “superstrict by tribe”: 1,049 genes, and “superstrict excluding hybrids”: 303 genes; Table 1). In the superstrict by tribe routine, putative paralogous genes were removed by tribe, meaning that a relatively high number of genes could be kept for further analyses (see STAR Methods for details and Methods S1C and S1D). In addition, we used both supermatrix maximum likelihood (ML) and coalescent-based approaches in ASTRAL-III⁴⁸ and ASTRAL-Pro⁴⁹ to infer the species phylogeny.

Node support was high for most tribes, regardless of the sampling routine or phylogenetic approach (Figure 2; Data S1E and S2), providing strong support for the monophyly of many (but not all) of the tribes. For example, from the supermatrix approach of the superstrict routine (concatenation of 297 target genes; total length 700,445 bp; 82.7% complete), the median support values across tribal nodes were 100% for bootstrap (BS), 50% for gene concordance factor (gCF), and 64% for site concordance factors (sCFs). From the coalescent-based approach (strict routine), the median local posterior probability (LPP) across tribal nodes was 100%, and support for the first quartet (Q1) was 78%. Tribes Camelineae and Iberideae were polyphyletic in all nuclear phylogenies (Data S2).

Based on the nuclear dataset, we consistently found tribe Aethionemeae sister to the rest of the Brassicaceae (hereafter defined as “core Brassicaceae”), main lineage III sister to a clade of lineages I–V, and lineages II + V sharing a common ancestor (Figures 1A and 1B).

Based on the superstrict routine (361 samples; Table 1), support for the split between Aethionemeae and the core Brassicaceae was high (BS: 100%; gCF: 70%; sCF: 63%; LPP: 100%; Q1: 70%; Figure 2; Data S2A), whereas support for the five remaining main lineages varied. Although main lineage III received good support (100%, 40%, 58%, 100%, and 91%, respectively), support for the remaining main lineages was relatively poor in terms of gCF (ranging from 1% to 5%) and sCF (ranging from 36% to 58%) values (Data S1F). Remarkably, BS and LPP values for these main lineages were (nearly) always 100%. The generally low support for the phylogenetic backbone (focusing on gCF, sCF, and Q1) was reflected in the topological differences among the family phylogenies based on the different sampling routines

(Figures 1A–1C) and was visualized as a large, complex reticulate core in our split network analysis (Figure 1D). Our approach to increase backbone support—and thereby confidence in the family’s main lineage relationships—with our superstrict excluding hybrids routine indeed resulted in higher support for all main lineages (gCF now ranging from 16% to 76%; range of sCF unchanged, but support for all main lineages the same or slightly higher; Data S1F). However, this came at the cost of removing many samples (and therefore species, genera, and tribes), with now only 138 samples left (Table 1).

Our inclusive routine resulted in placement of main lineage I as sister to a clade of main lineages II + IV + V (Figure 1A), whereas the stricter routines of the nuclear dataset showed a consistent placement of lineage IV as sister to a clade of lineages I + II + V (Figure 1B) in both the ML and coalescent-based approaches. This showed that removing a first set of most variable genes (strict routine) considerably impacts the backbone of the topology, whereas an additional removal of less variable genes (superstrict, superstrict by tribe, and superstrict excluding hybrids) has little effect.

We used Townsend’s phylogenetic informativeness⁵⁰ to quantify the impact of each of the 1,081 nuclear genes on the final species tree (Figure S1). Gene informativeness varied markedly, with genes from the B764 bait set on average more informative than those from the A353 bait set, which likely reflects the family-specific design of the B764 bait set.

Extended global plastome Brassicaceae phylogeny

Our plastome family ingroup sampling included 502 samples, covering 438 species, 266 genera (76%), and all 58 tribes (following German et al.²⁴), with raw input data from newly sequenced samples (Data S1A) and previously published data from Nikolov et al.³ (Data S1B) and Walden et al.² (Data S1G).

Similar to the nuclear phylogeny, tribal support was generally high, with median nodal sCF of 81% and BS of 100% (Figure 3; Data S1F). As in the nuclear phylogeny, tribe Aethionemeae was sister to all remaining Brassicaceae. Tribes within lineage II were similar to those in the nuclear phylogeny. However, cytonuclear discordance manifested by among others the following topological differences (Figure 4): (1) in the plastome phylogeny, lineage I (not III) was sister to all other lineages (II–V), with lineage III sister to a clade formed by II + IV + V; (2) lineage II was polyphyletic in the plastome phylogeny, with several tribes (Aphragmeae, Coluteocarpeae, Conringieae, Kernereae, and Plagiolobeae) recovered within lineage V; (3) tribe Stevenieae, recovered within lineage IV in the nuclear phylogeny, was recovered within lineage I in the plastome phylogeny; and (4) new tribe Asperuginioideae (see below), assigned to lineage IV in the nuclear phylogeny, formed a clade with Biscutelleae in the plastome phylogeny, and these tribes together were sister to all other remaining lineages (II + IV + V).

(C) Cladogram of main lineages as recovered from plastome supermatrix ML approach with its node support (BS/sCF). The † indicates a major polyphyly in main lineage II (supertribe Brassicodae).

(D) Split network of the mustard family tribes computed from uncorrected p-distances on a supermatrix of the nuclear genes covered by the superstrict routine, covering 317 genera of 56 tribes. The network highlights the complex reticulate evolution both in the ancestors of extant main lineages, as well as within some of the main lineages. Names in bold highlight rogue taxa described in the main text. Inset drawings show the high morphological diversity within the family, which contains growth forms such as herbaceous, frutescent, and woody species, the latter comprising lianas, shrubs, and cushion plants. Drawings by Esmée Winkel, Naturalis Biodiversity Center.

Table 1. Sampling routine definitions

| Sampling routine | Inclusive | Strict | Superstrict | Superstrict by tribe | Superstrict excl. hybrids |
|--|------------------------|--------------|-------------|----------------------------------|---------------------------|
| Gene and sample selection criteria | | | | | |
| Gene minimum proportion of samples recovered | 0.1 | 0.2 | 0.2 | 0.2 | 0.2 |
| Gene minimum proportion of target length recovered | 0.1 | 0.2 | 0.2 | 0.2 | 0.2 |
| Sample minimum proportion of total target length recovered | 0.0 | 0.4 | 0.4 | 0.4 | 0.4 |
| Sample minimum proportion of genes recovered | 0.0 | 0.2 | 0.2 | 0.2 | 0.2 |
| Gene SNPs proportion threshold for all samples | none | outliers | 0.02 | 0.02 | 0.02 |
| Remove outlier genes per sample? | yes | yes | yes | yes | yes |
| Remove putative hybrids and rogue taxa? | no | no | no | no | yes ^a |
| Dataset summary | | | | | |
| Number of genes that passed HybPhaser criteria | | | | | |
| All genes | 1,081 (100) | 1,057 (97.8) | 306 (28.3) | 1,076 (99.5) | 306 (28.3) |
| B764 genes | 728 ^b (100) | 721 (99) | 194 (26.6) | 728 (100) | 194 (26.6) |
| A353 genes | 353 (100) | 336 (95.2) | 112 (31.7) | 348 (98.6) | 112 (31.7) |
| Number of genes that passed de-noising loop and used in phylogenetic analysis | | | | | |
| All genes | 1,018 (94.2) | 1,013 (93.7) | 297 (27.5) | 1,049 (97) (1,031 ^c) | 303 (28) |
| B764 genes | 684 (94) | 688 (94.5) | 189 (26.0) | 709 (97.4) | 193 (26.5) |
| A353 genes | 334 (94.6) | 325 (92.0) | 108 (30.6) | 340 (96.3) | 110 (31.2) |
| Number of samples in final dataset | 380 | 361 | 361 | 361 | 138 |
| Number of species | 375 | 356 | 356 | 356 | 138 |
| Number of genera as currently accepted | 319 (332) | 305 (317) | 305 (317) | 305 (317) | 124 |
| Number of tribes ^d | 57 | 56 | 56 | 56 | 36 |
| Coalescent-based results from ASTRAL-III | | | | | |
| LPP (mean of all nodes) | 0.94 | 0.95 | 0.92 | 0.88 | – |
| LPP (median of all nodes) | 1 | 1 | 1 | 1 | – |
| First quartile proportion (mean of all nodes) | 0.55 | 0.56 | 0.57 | 0.56 | – |
| First quartile proportion (median of all nodes) | 0.48 | 0.50 | 0.50 | 0.49 | – |

Definition of sampling routines and criteria used in HybPhaser to select nuclear genes for downstream analyses, with dataset summary and coalescent-based results from ASTRAL-III. Values in parentheses are percentages relative to the total number of genes in each bait set (728 for B764, i.e., 764 minus 36 overlapping genes; 353 for A353). See also [Methods S1](#).

^aPutative hybrids were selected based on locus heterozygosity and allele divergence, with all samples within the upper 50% removed; rogue tribes as discussed in the main text.

^bThis corresponds to the 764 genes from the bait set minus 36 genes overlapping with the A353 bait set.

^cMean number across samples, with genes retained differently in different tribes (i.e., tribe-specific). As a result, more genes could be retained in total as compared with the inclusive routine, but at a lower mean sample occupancy across all genes, thus leading to a relatively sparse matrix.

^dTribe definitions following latest taxonomic insights presented in German et al.,²⁴ i.e., 58 tribes in total.

Topological incongruences within tribes between the nuclear and plastome phylogenies could be calculated for 23 tribes for which the number of shared species was >3 ([Data S1H](#)). Results varied from full concordance (tribes Biscutelleae, Chorisporeae, Cremolobeae, Descurainieae, Eudemaeae, Heliophleae, Isatideae, and Physarieae; generalized Robinson-Foulds distance 0) to full discordance (tribe Lepidieae only; generalized Robinson-Foulds distance 1), with a median generalized Robinson-Foulds distance across all tribes of 0.31.

Allelic variability highlights polyploid origins

We assessed allelic variation using HybPhaser v2.0⁴⁷ by calculating the proportions of loci with divergent alleles (% locus heterozygosity [LH]) and average divergence between alleles (% allele divergence [AD]), using the 1,013 nuclear genes included in the strict routine (ignoring 38 samples with coverage too low

to calculate LH and AD; [Data S1E](#)). Although most samples fell into the expected range of “normal” diploid species (roughly LH < 90% and AD < 1), a number of samples showed signs of polyploidization ([Figures 5](#) and [S2](#)). Mean values for both LH and AD were high at 75.1% and 2.25%, respectively, with no clear difference between genes from the B764 and A353 bait sets ([Data S1E](#) and [S1I](#)).

Based on the list of LH and AD values for all species, we tentatively distinguished four family-specific classes: “hybrid,” “old polyploid,” “highly polyploid,” and “old and highly polyploid” ([Figure 5](#); see [STAR Methods](#) for details and class circumscription). Nearly all representatives of tribe Thelypodieae were highly polyploid, suggesting that the tribe experienced one or more recent polyploidization events. Similarly, all representatives of tribe Lepidieae fell within the hybrid or highly polyploid classes, and also, tribes Alysseae, Brassiceae, Cardamineae,

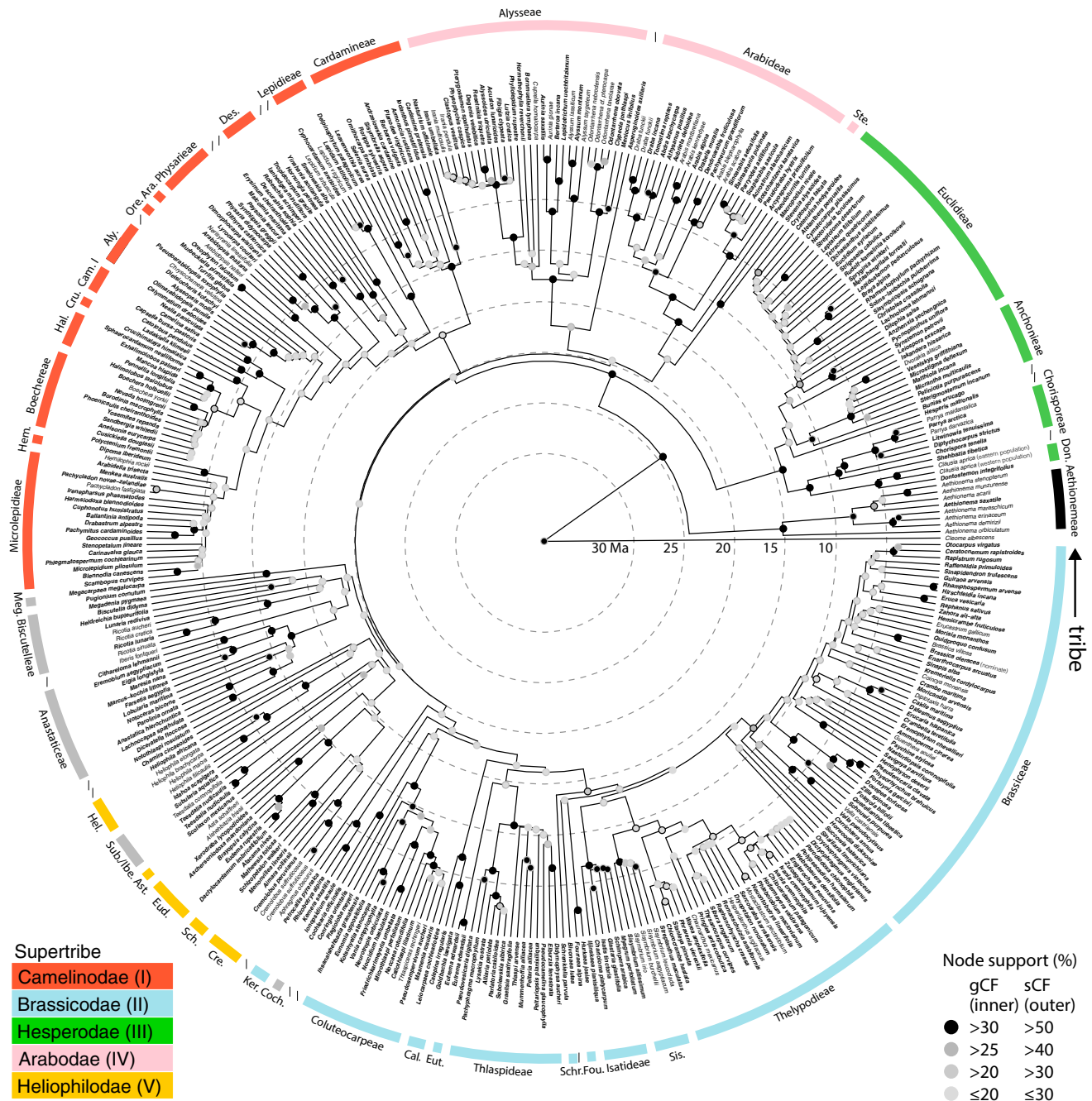


Figure 2. Time-calibrated genus-level Brassicaceae Tree of Life (BrassiToL) from a maximum likelihood analysis of a 297 nuclear genes supermatrix (superstrict routine)

Genus type species are highlighted in bold. All tribes with more than a single representative are listed. Abbreviations of tribes are as follows: Aly., Alyssoisideae; Ara., Arabidopsidae trib. nov.; Ast., Asteae; Cal., Calepineae; Cam. I., Camelineae I; Coch., Cochlearieae; Cre., Cremolobae; Cru., Crucihimalyae; Des., Descurainieae; Don., Dontostemoneae; Eud., Eudemeae; Eut., Eutremeae; Fou., Fourraeae; Hal., Halimolobae; Hel., Heliophleae; Hem., Hemilophleae; Ibe., Iberideae; Ker., Kernereae; Meg., Megacarpaeae; Ore., Oreophytoneae; Sch., Schizopetaleae; Schr., Schrenkielleae trib. nov.; Sis., Sisymbrieae; Ste., Stevenieae; Sub., Subularieae. See also [Data S2A](#) for a fully annotated version of this phylogeny (including bootstrap, gCF, sCF and node age 95% HPD intervals) with outgroups representing all families within the order Brassicales and calibration nodes. See also [Data S2A](#).

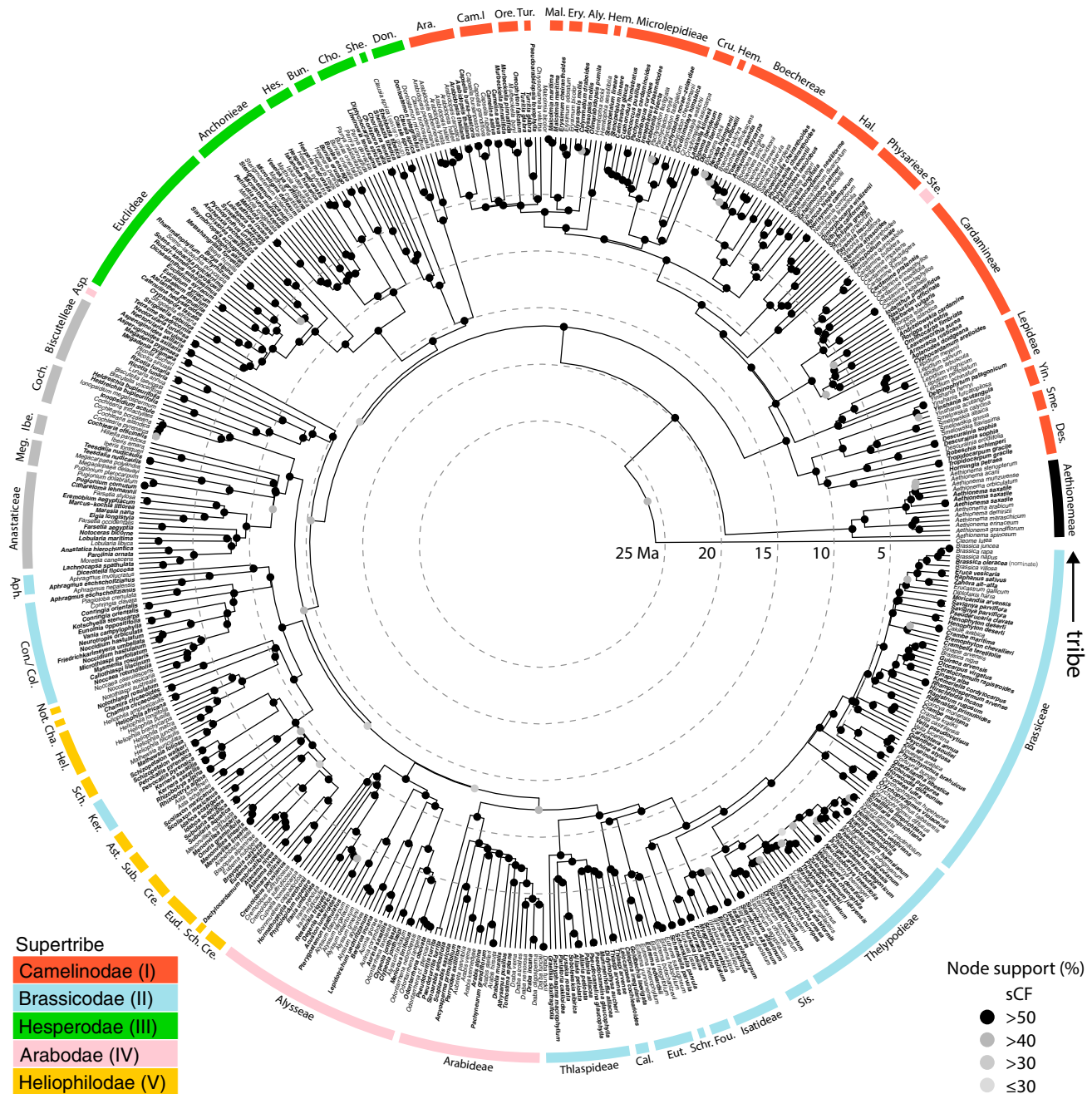


Figure 3. Time-calibrated genus-level Brassicaceae Tree of Life from a maximum likelihood analysis of a 60 plastome genes supermatrix
Genus type species are highlighted in bold. All tribes with more than a single representative are listed. Abbreviations of tribes follow those of Figure 2, with additionally: Aph., Aphragmeae; Asp., Asperuginoideae trib. nov.; Bun., Buniadeae; Cha., Chamireae; Cho., Chorisporae; Col., Coluteocarpeae; Con., Conringieae; Don., Dontostemoneae; Ery., Erysimeae; Hes., Hesperideae; Mal., Malcolmieae; Not., Notothlaspidiae; Ore., Oreophytoneae; She., Shehbazieae; Sme., Smelowskieae; Tur., Turritidae; Yin., Yinshanieae. See also Data S2B for a fully annotated version of this phylogeny (including bootstrap, sCF and node age 95% HPD intervals) with outgroups representing all families within the order Brassicales and calibration nodes. See also Data S1.

and Microlepidieae had the bulk of their representatives in one of the polyploid classes (Figure 5).

Our phylogenetic results highlighted several jumpy or rogue taxa (mostly tribes) that were recovered in different positions in the phylogenies from different sampling routines. Most importantly, in the nuclear supermatrix ML approach and the stricter

coalescent-based approaches (Figures S3A and S3D–S3F), Anastaticeae, Biscutelleae, and Megacarpaeae formed a clade sister to lineage V, whereas in the inclusive coalescent-based approaches (ASTRAL-III and ASTRAL-Pro; Figures S3B and S3C), Megacarpaeae was sister to Anastaticeae, with the two tribes sister to lineage II. The position of tribe Biscutelleae

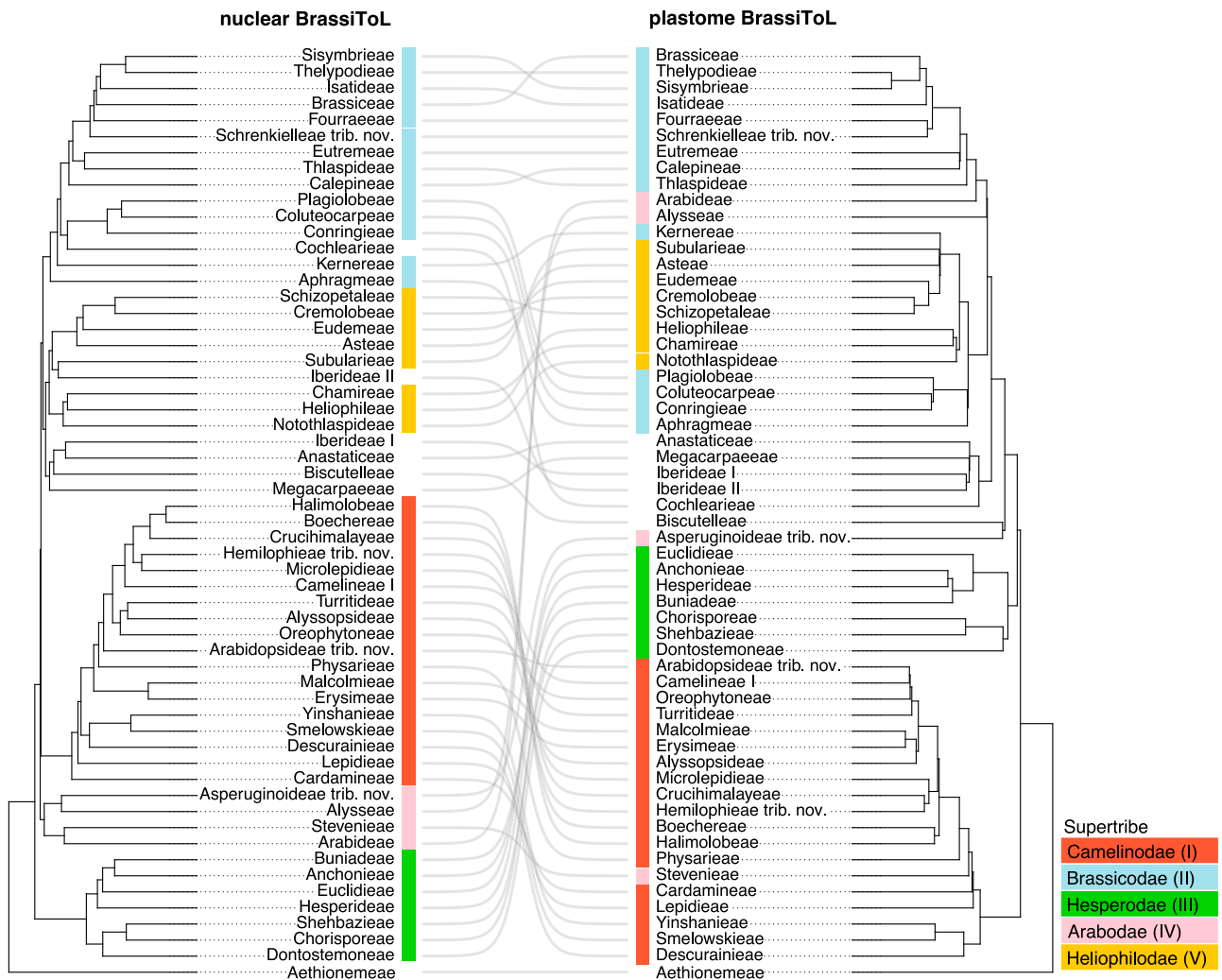


Figure 4. Cytonuclear discordance at tribe level in newly derived nuclear and plastome Brassicaceae Trees of Life

Curved lines between tip labels from the two phylogenies link the same tribes. Tribes are represented by a randomly chosen sample from each tribe. Rogue tribes have not yet been assigned to a supertribe due to their changing topological position from different sampling routines and phylogenetic approaches.

changed depending on the coalescent-based approach used (ASTRAL-III vs. ASTRAL-Pro). Contrary to our plastome phylogeny, we recovered genus *Iberis* (tribe Iberideae) as sister to tribe Anastatiaceae in our nuclear phylogenies, and not as sister to *Teesdalia*, the other genus assigned to tribe Iberideae. Species that belong to one of the rogue taxa generally also show relatively high LH and AD values (17 of the 29 samples fall within the “realm of polyploids,” with another 6 samples bordering it; Figure 5), suggesting a polyploid origin.

Fossil calibration shows an Eocene origin of the mustard family

We used the Turonian *Dressiantha bicarpellata* fossil (93.6–89.3 Ma⁵¹) to calibrate the stem node of order Brassicales (Data S2A and S2B). We validated our dating estimates with nine other fossils and biogeographical events (Data S1J), as suggested by Franzke et al.⁵² In general, results from our nuclear phylogeny supported the expected dates (based on literature) or were

younger. In the plastome phylogeny, our age estimates were significantly younger. We found good corroboration with age estimates for the Miocene *Cappariodoxylon holleisii* fossil (16.3 Ma^{53,54}) in the nuclear phylogeny (poor in the plastome phylogeny), the maximum crown age of the Mediterranean genus *Ricotia* (11.3–9.2 Ma⁵⁵) in both phylogenies, the estimated age of the most recent common ancestor of the *Arabis alpina* clade (3.27–2.65 Ma⁵⁶) in the nuclear phylogeny (poor in the plastome phylogeny), and the maximum age of the New Zealand alpine genus *Pachycladon* (max. 1.9 Ma^{57,58}) in both phylogenies. Support was medium in the nuclear phylogeny (and poor in the plastome phylogeny) for the Paleocene fossil *Akania* sp. (~61 Ma^{59,60}) and the early Tertiary *Palaeocleome lakensis* fossil (55.8–48.6 Ma⁶¹). Support was poor for the vicariance event in *Clausia aprica*⁶² and the maximum age of Hawaiian *Lepidium*,⁶³ with recovered ages in both the nuclear and plastome phylogenies actually older than expected. Corroboration with the Brassicaceae *Thlaspi primaevum* fossil (~32 Ma^{64–66}) was poor

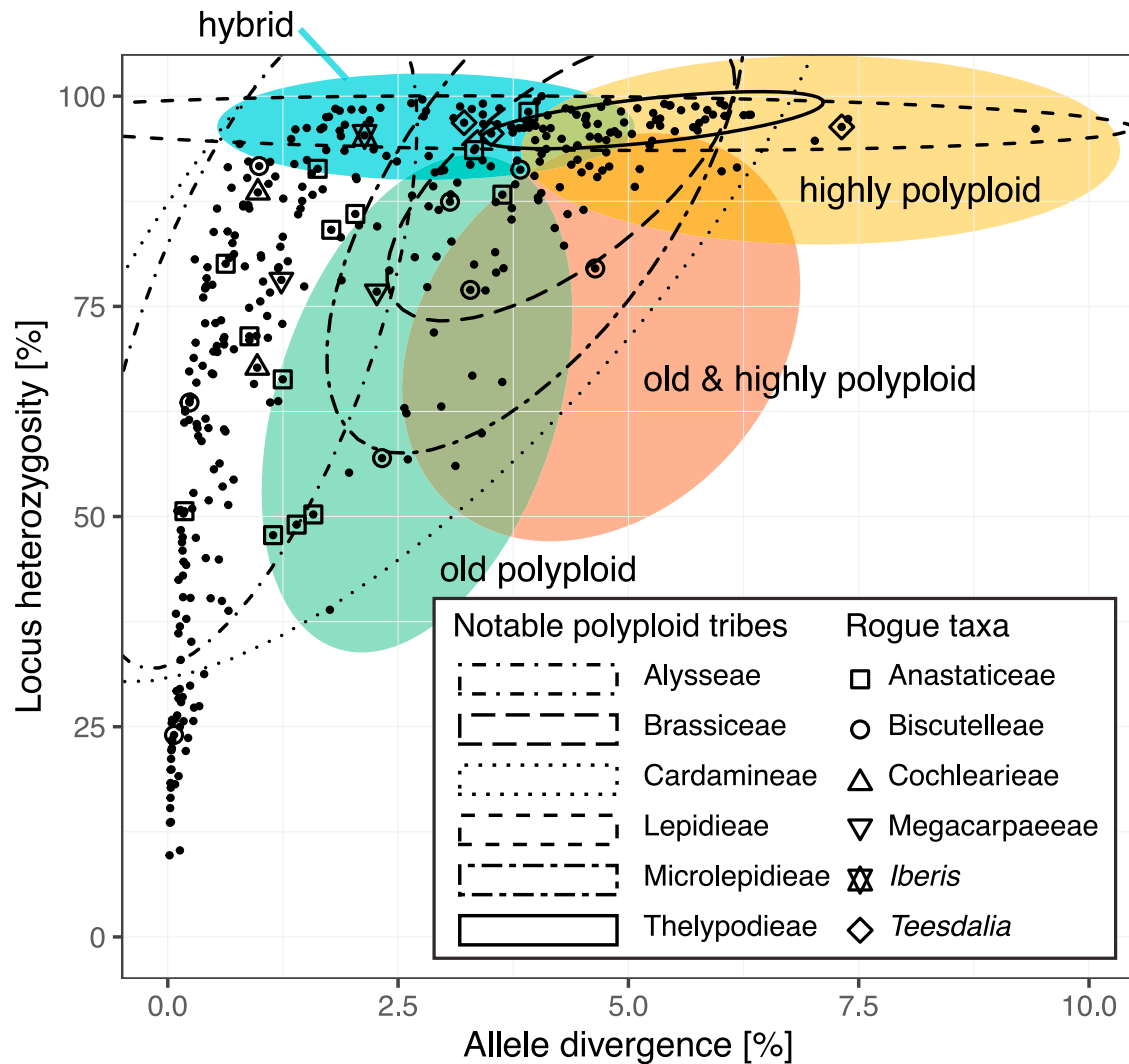


Figure 5. Scatterplot displaying the locus heterozygosity and allele divergence of all samples included in the strict routine

Colored ovals indicate four classes in the expected realm of polyploids, rough estimates of what could be considered likely hybrids, polyploids, and their ages (see main text for the rationale behind assigning these different classes). The overlap between these classes highlights the uncertainty. The six largest tribes (represented by >10 samples) for which (most) of their samples fall within this realm of polyploids are annotated using ellipses showing the 95% confidence level for a multivariate t-distribution of their data. Samples from rogue taxa are highlighted using different shapes and colors. For more details, see [Data S1E](#) and [S1I](#) and [Figure S2](#).

(node estimated at 10.8 Ma) in both phylogenies, which is unsurprising, given the ambiguous taxonomic identity of this fossil.⁵²

Based on our nuclear dataset, we found that the core Brassicales (Brassicaceae + Cleomaceae) and Capparaceae split around 43.2 Ma (95% highest posterior density, or HPD: 45.0–41.1) in the middle Eocene ([Data S2A](#)). Subsequently, the Brassicaceae split from sister family Cleomaceae around 38.8 Ma (95% HPD: 40.5–36.9) in the middle to late Eocene ([Data S1J](#)). The core Brassicaceae split from tribe Aethionemeae around 24.5 Ma (95% HPD: 25.7–23.1), in the late Oligocene. All five main lineages (supertribes) in the core Brassicaceae originated during rapid radiation in the early to middle Miocene (median stem ages 21.2–19.8 Ma; median crown ages 19.9–14.4 Ma; [Figure S4](#); [Data S1F](#)). Mean median stem age across tribes was 12.1 Ma (max. 18.9 Ma, Subularieae; min. 4.0 Ma, Boechereae;

$n = 52$), whereas mean median crown age was 8.0 Ma (max. 18.1 Ma, Subularieae; min. 0.2 Ma, Oreophytoneae; $n = 37$). Results from our plastome dataset were much younger than those from our nuclear dataset, with a family crown age of 20.2 Ma (95% HPD: 29.0–13.0) and a core Brassicaceae crown age of 16.9 Ma (95% HPD: 24.3–10.2; [Data S1F](#)).

DISCUSSION

Unprecedented genus-level BrassiToL

We present here the most complete nuclear and plastome-derived BrassiToL to date, together based on more than 1,000 genes and covering 92% of the 349 genera, representing all 58 tribes ([Figures 1, 2, and 3](#); [Data S2](#)). This global genus-level BrassiToL will be an indispensable tool for family experts

interested in understanding the evolutionary processes that shaped current biodiversity patterns. In addition, many scientists working on *Arabidopsis* or other Brassicaceae model species or crops will benefit from this improved phylogenetic framework when investigating gene regulatory mechanisms across the family or exploring specific traits related to stress/disease in crops and their wild relatives.

The backbone of our nuclear BrassiToL largely agrees with that of Nikolov et al.³ (based solely on the Brassicaceae-specific bait set targeting 764 genes), which included 63 species covering most tribes. Specifically, our nuclear BrassiToL confirms (1) the recognition of tribe Aethionemeae as sister to the rest of the Brassicaceae (i.e., core Brassicaceae) and (2) main lineage III (supertribe Hesperodae, Figures 1A and 1B) as sister to the remaining core Brassicaceae.^{3,33,34} The relationships among the other main lineages varied depending on the applied sampling routine and/or phylogenetic approach, although the sister relationship between main lineages II (Brassicodae) and V (Heliophilodae) was consistently recovered as well (Figures 1A and 1B). We are confident that our expanded dataset and more diverse analyses of putative single-copy markers within Brassicaceae offer the most trustworthy estimation of the family relationships to date. We acknowledge, however, that—even with this massive dataset and careful analyses—we have not fully resolved the well-known problem of recovering deeper nodes in the Brassicaceae phylogeny (see below).

When comparing our plastome phylogeny (representing 265 currently accepted genera, all 58 tribes) with the nuclear phylogeny (representing 319 currently accepted genera, 57 tribes), we found strong cytonuclear discordance among the main lineages, as recently also demonstrated by Nikolov et al.³ and Mabry et al.³³ (Figure 4). At a lower taxonomic level, such cytonuclear discordance was studied in more detail for *Arabidopsis* and its putative close relatives,⁴¹ as well as tribe Biscutelleae,⁴³ where it was hypothesized that complex hybridization and introgression events caused topological incongruences. For example, tribe Biscutelleae harbors four genus-specific WGDs resulting from hybridization between parental genomes belonging to the same lineages, closely related lineages, and even two different supertribes⁴³ (Camelinodae and Brassicodae).

Toward resolving the family phylogeny backbone

Relationships at the shallower nodes are generally well-resolved in our phylogenies, leading to strongly supported relationships among genera within most of the tribes (e.g., Q1 node support was >50% in 34 of the 42 tribes for which we could calculate node support, i.e., tribes represented by >1 species; Figure S4; Data S1F). Interestingly, most of the deepest nodes (connecting the Brassicales families) are generally also well supported (Figure S4). However, the deeper nodes *within* the Brassicaceae family—reflecting the relative positions of the five supertribes in the subfamily Brassicoideae—have been notoriously hard to recover in the past and remain incompletely resolved even with more than a thousand genes to study (Figure 1). Importantly, we show that high BS support (used in the past to claim a solid family backbone³⁵) can be recovered nonetheless, even for conflicting topologies from different sampling routines, thereby challenging the value of BSs in phylogenomics. Instead, metrics on node support that take underlying gene variation into account

(gCF; Q1, first quartet from coalescent-based analysis) and single-nucleotide polymorphisms (SNPs) (sCF) are more informative when assessing branch support from genomic datasets.⁶⁷

Low backbone support in a phylogenetic tree can have several causes: (1) biological processes within the family, such as gene duplication and loss, hybridization, incomplete lineage sorting, a rapid radiation, and gene saturation, (2) one or more artifacts (e.g., deficient or erroneous data such as paralogs interpreted as orthologs), or (3) a combination thereof.⁶⁸ As gene saturation is a process that increases over time, we believe that it is not influencing our dataset to a large extent because node support among Brassicales families is generally higher than support for main lineages (subfamilies and supertribes) within Brassicaceae (Figure S4). Our dataset clearly indicates, however, that hybridization events have occurred frequently throughout Brassicaceae evolution. We found LH and AD to be very high in many species, with mean values of 75.1% and 2.25%, respectively (Data S1E and S1I). In comparison, these values were only 52.3% and 0.21%, respectively, across all non-hybrid natural accessions of pitcher plants (*Nepenthes* spp.) and 89.4% and 0.86%, respectively, in their known and suspected hybrids.⁴⁷ This highlights the abundant presence of hybrids in Brassicaceae, such as in tribes Alysseae, Brassiceae, Cardamineae, Lepideae, Microlepidaeae, and Thelypodieae, and rogue tribes Anastaticae, Biscutelleae, Cochlearieae, Iberideae, and Megacarpaeae (Figure 5; see next section).

Brassicaceae are notoriously known for their rampant hybridization (see below), and it is challenging to include all such evolutionary oddballs into a single evolutionary model to recover the family's “true” systematic relationships. A network approach could potentially represent these relationships more faithfully⁶⁹ (Figure 1D). Aware of the issues with data deficiency and paralogs, we specifically designed our analyses to disentangle some of the deeper nodes by stepwise removal of putative paralogous genes and samples of a putative hybrid origin (“inclusive,” “strict,” “superstrict,” “superstrict by tribe,” and “superstrict by tribe excluding hybrids” routines; Table 1; see STAR Methods for details). A recent alternative approach to exclude paralogs using synteny between whole-genome sequences⁷⁰ showed similar differences between tree topologies at the deeper nodes as those among our inclusive and stricter datasets (Figures 1A and 1B). Specifically, the branching order of clades I–IV based exclusively on strict syntenic orthologs was identical to the branching order recovered in our stricter datasets (Figure 1A), lending independent support for the importance of our substantial paralog filtering routines.

Interestingly, we found that going from the strict routine (removing only outlier genes and keeping 97.8% of all genes; Table 1) toward the superstrict routine (removing all genes with a mean SNP proportion >0.02, keeping only 28.3% of all genes), the topology of our BrassiToL from the coalescent-based approach hardly changed (Figure S3). We therefore believe that the superstrict routine (Figure 2) provides the best estimate of the phylogeny, as it includes most samples and at the same time excludes any “unnecessary” genes.

Rogue taxa share a mesopolyploid origin

Our results corroborate with previous studies that highlighted several difficult-to-place or rogue taxa,³ including tribes

Anastatiaceae, Biscutelleae, Cochlearieae, and Megacarpaeae, and the genera *Iberis*, *Idahoa*, and *Subularia*. Contrary to the bulk of the tribes, which were consistently placed within a main lineage in our different nuclear phylogenies, these taxa “jumped” positions across the different phylogenies (Figure S3). This has previously been interpreted as the result of inter-tribal or inter-lineage hybridizations between distantly related parental species (genus- or tribe-specific meso-polyploidizations), as explained in the next paragraph.

The genus *Brassica* and the tribe Brassiceae were the first Brassicaceae taxa reported with a WGD postdating the family-specific palaeotetraploidization At- α event.^{71–73} Since this pioneering work, more than a dozen genus- and tribe-specific mesopolyploid WGDs have been discovered throughout the family.^{9,39,43,74} Because most mesopolyploid taxa have an allopolyploid origin arising from hybridization between distantly related parental species,^{42,43,75} inferring their phylogenetic position within the family tree is often a challenging endeavor. For all the rogue taxa we identified, a mesopolyploid origin has indeed been demonstrated or claimed. For instance, for tribe Cochlearieae, whole-genome triplication has been demonstrated, whereas Anastatiaceae and *Iberis* have a mesotetraploid origin.³⁹ Biscutelleae harbor four different WGDs specific to *Biscutella*, *Heldreichia*, *Lunaria*, and *Ricotia*.^{43,76} Both genera of Megacarpaeae (*Megacarpaea* and *Pugionium*) have been shown to be formed by independent WGDs.^{77,78} *Teesdalia*, based on available chromosome numbers ($2n = 36$ in *T. coronopifolia* and *T. nudicaulis*, $2n = 20$ in *T. conferta*), most likely has a polyploid origin, but (cyto)genomic data are lacking. Interestingly, based on our results, not all mesopolyploids act as rogue taxa, including tribe Brassiceae.

Brassicaceae originated during Earth’s icehouse era

Our results suggest a somewhat younger age for the Brassicaceae than previously published (Figure 2; Table 2). We recovered a middle to late Eocene stem age for the mustard family (38.8 Ma; 95% HPD: 40.5–36.9), with a late Oligocene crown age (24.5 Ma; 95% HPD: 25.7–23.1). Although the family’s crown node age estimates ranged from 54.3¹⁹ to 15.0 Ma¹⁸ in the earlier studies, results from more recent Brassicaceae studies converged to a crown age of 37.1–32.4 Ma, which appears insensitive to data type (nuclear/plastome), methods, and fossils used.²³

Genomic datasets like ours are notoriously difficult to use in time calibration due to the sheer amount of data that needs to be analyzed simultaneously.⁸⁵ There is no consensus about the best approach to retrieve reliable dating estimates based on big genomic datasets, but our experience is that including a subset of only clock-like genes yields a more reliable time-calibrated tree compared with a more inclusive dataset (including, e.g., all genes), which commonly results in a forward shift in time.

Based on our time-calibrated analyses using the 20 most clock-like genes, the family’s origin and the onset of its diversification coincide with the cooling of the Earth during the Eocene-Oligocene transition (so-called greenhouse to icehouse transition^{86,87}). This period was characterized by a worldwide replacement of tropical forests with temperate forests, open vegetation, and deserts, which are all typical habitats of extant Brassicaceae. Tribe Aethionemeae and the five supertribes in subfamily Brassicoideae originated quickly after in the early

Miocene (median stem ages range from 21.2 to 19.8 Ma; median crown ages range from 19.9 to 14.4 Ma; Data S1F). A combination of short branch lengths and low support in the BrassiToL around these events (Figure S4) supports the idea of an early rapid radiation of the family.

Results from our plastome phylogeny generally show a ~5 Ma forward shift in time relative to our nuclear phylogeny, possibly the result of including all 60 plastome genes in our study, compared with a subset of 20 clock-like genes in our nuclear analysis (Figure 3; Table 2). Median crown age of the family was 20.2 Ma (vs. 24.5 Ma in nuclear analysis), and median crown age of core Brassicaceae 16.9 Ma (vs. 21.1 Ma). Results from our plastome study are also much younger (nearly 10 Ma) than previously found by Walden et al.² based on partly the same dataset (but a different set of fossils used for calibration; Table 2).

Improved Brassicaceae phylogeny warrants updated family classification

Our study provides new systematic insights and consolidates results from recent phylogenetic studies in Brassicaceae, fully supporting the new family classification of German et al.²⁴ This includes the formal definition of two subfamilies (Aethionemoideae, including only *Aethionema*, and Brassicoideae), five supertribes (Camelinodae, previously lineage I; Brassicodae, lineage II; Hesperodae, lineage III; Arabodae, lineage IV; and Heliophilodae, lineage V), and 58 tribes.

At the tribal level, our results confirm that the scientifically important genus *Arabidopsis* is not closely related to any other genera traditionally placed in tribe Camelinaeae,^{3,22} supporting the movement of *Arabidopsis* into a new monogeneric tribe, Arabidopsidae (supertribe Camelinodae²⁴). Our results also support the following taxonomic changes as recently published by German et al.²⁴: (1) erecting the monospecific and distinct genus *Asperuginoides* to its own new tribe, Asperuginoidae, (2) combining the genera *Dipoma* and *Hemilophia* in the new tribe Hemilophiae, (3) combining the genera *Idahoa* and *Subularia* in the re-established tribe Subularieae (plastome phylogeny only), and (4) raising the monospecific and distinct genus *Schrenkiella* into its own tribe, Schrenkielleae. We found all tribes to be monophyletic and highly supported in both nuclear and plastome phylogenies, except for Camelinaeae (polyphyletic in both nuclear and plastome phylogenies), Iberideae (polyphyletic in the nuclear phylogeny), and Subularieae (polyphyletic in the nuclear phylogeny). These tribes, in addition to the aforementioned rogue and hybrid tribes, require further systematic studies to uncover their exact evolutionary history and determine their taxonomic status.

Conclusions

We provide the first global, calibrated, nuclear, and plastome BrassiToL, offering an important step forward in untangling the notoriously difficult phylogenetic relationships across Brassicaceae. We applied the latest bioinformatic tools to select the most reliable genes to construct the species tree, thereby increasing topological accuracy and highlighting likely polyploid taxa. Our improved phylogenetic framework supports the reinstatement of the new family classification by German et al.,²⁴ including two subfamilies (Aethionemoideae and Brassicoideae),

Table 2. Brassicaceae divergence time estimates

| Studies | Crown Brassicaceae (Ma) | Crown core Brassicaceae ^a (Ma) | Method | Dataset | Calibration |
|--|-------------------------|---|---|---|--|
| Koch et al. ⁷⁹ | – | 25.9–23.1 | synonymous substitution rate | <i>Adh</i> and <i>Chs</i> | synonymous substitution rate |
| Franzke et al. ¹⁸ | 35.0–15.0–1.0 | 28.0–11.0–1.0 | BEAST | <i>nad4</i> | one secondary calibration |
| Beilstein et al. ¹⁹ | 64.2–54.3–45.2 | 54.3–46.9–39.4 | BEAST | <i>ndhF</i> and <i>PHYA</i> | four fossils |
| Couvreux et al. ²⁰ | 49.4–37.6–24.2 | 43.8–32.3–20.9 | BEAST | eight genes from nuclei, chloroplast, and mitochondria | one fossil |
| Kagale et al. ⁸⁰ | – | 26.6 | synonymous substitution rate | 213 nuclear orthologs | synonymous substitution rate |
| Edger et al. ¹¹ | 45.9–31.8–16.8 | – | BEAST | 1,115 single-copy nuclear genes | two fossils |
| Hohmann et al. ⁹ | 38.6–32.4–27.1 | 27.3–23.4–19.9 | BEAST | plastomes | four fossils |
| Huang et al. ²² | 37.8–37.1–36.3 | 30.3–29.7–29.1 | r8s | 113 low-copy nuclear orthologs | 18 fossils ^b |
| Cardinal-McTeague et al. ⁵⁴ | 44.1–37.7–31.4 | – | BEAST | Chloroplast DNA (<i>ndhF</i> , <i>matK</i> , <i>rbcL</i>) and mitochondrial DNA (<i>matR</i> , <i>rps3</i>) | three fossils ^b |
| Mohammadin et al. ⁸¹ | 58.9–48.0–37.5 | 35.4 | BEAST | plastomes | one secondary calibration |
| Guo et al. ⁸² | 41.8–34.9–29.0 | 29.8–25.1–21.3 | MCMCTree | plastomes | 14 fossils ^c |
| Mandáková et al. ³⁹ | 54.7–40.1–29.4 | 30.6 | BEAST | plastomes | four fossils |
| Huang et al. ²³ | 33.2–29.9–26.8 | 22.9–21.3–19.6 | BEAST | plastomes | four fossils |
| Ramírez-Barahona et al. ⁸³ | 52.7–41.9–30.5 | – | BEAST | <i>rbcL</i> , <i>atpB</i> , <i>matK</i> , <i>ndhF</i> , 18S, 26S, 5.8S | 238 fossils, angiosperm-wide |
| Walden et al. ² | 35.7–29.9–24.3 | 29.6–25.1–20.9 | BEAST | plastomes | four fossils |
| Legalov et al. ⁸⁴ | – | >36.4 | indirect calibration via phytophagous beetles | – | <i>Ceutorhynchus</i> beetle fossils |
| This study (nuclear dataset) | 25.7–24.5–23.1 | 22.4–21.1–19.9 | treePL | phylogeny using superstrict routine; calibration using subset of 20 most clock-like genes | one fossil validated by nine secondary calibration points |
| This study (plastome dataset) | 29.0–20.2–13.0 | 24.3–16.9–10.2 | treePL | plastomes | one fossil validated by eight secondary calibration points |

Comparison of estimates from past studies and the current study (taken and updated from Huang et al.²³).

^aCore Brassicaceae are all Brassicaceae, excluding basal tribe Aethionemeae.

^bDating results excluding *Thlaspi primaevum*.

^cOnly results that exclude Brassicales fossils.

with five supertribes in the latter (Arabodae, Brassicodae, Camelinodae, Heliophilodae, and Hesperodae), and a total of 58 tribes of which five newly described or re-established.

The ultimate goal of our Brassicaceae consortium is to build a complete Brassicaceae family phylogeny including all ~4,000 species. The methods that we applied (including our “mixed baits” target capture sequencing from—sometimes over 200 years old—herbarium material⁴⁶) have proven successful and are becoming more affordable (now roughly € 40 per sample) and can easily be scaled up. Such a new species-level phylogeny will further boost the significance of this model plant family—for both fundamental and applied research programs covering all fields of plant biology.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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● **QUANTIFICATION AND STATISTICAL ANALYSIS**

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cub.2023.08.026>.

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AUTHOR CONTRIBUTIONS

F.L., K.M., and K.P.H. designed the study. F.L., K.M., K.P.H., I.A.A.-S., C.D.B., C.K., A.H.v.H., D.A.G., M.A.K., Ó.T.-N., B.Ö., V.R.I., O.M., N.M.H., M.T., M.D.W., I.R., S.Š., B.N., R.V., C.B., H.R., S.B.J., M.S., A.F., A.G., S.Z., B.J.L., N.S., F.W.S., I.S., P.H., W.J.B., and F.F. contributed to the sampling process. K.P.H., A.H.v.H., C.K., N.W., C.D.B., and A.R.Z. performed laboratory work. K.P.H., C.K., L.N., N.M.H., A.R.Z., and E.L. performed the various analyses. K.P.H., F.L., C.K., K.M., W.J.B., I.A.A.-S., C.D.B., A.H.v.H., L.N., D.A.G., M.A.K., M.A.L., and N.W. took the lead in writing the manuscript, with subsequent input from all co-authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---|--|---|
| Biological Samples | | |
| See Data S1A , S1B, and S1G | This study | N/A |
| Critical Commercial Assays | | |
| DNeasy PowerPlant Pro Kit | Qiagen, Hilden, Germany | Cat#13400 |
| DNeasy PowerClean Pro Cleanup Kit | Qiagen, Hilden, Germany | Cat#12997 |
| NEBNext Ultra II FS Kit | New England Biolabs, Ipswich, Massachusetts, USA | Cat#E7645L |
| IDT 10 index | Integrated DNA Technologies, Coralville, Iowa, USA | Cat#10008052 |
| NovaSeq 6000 sequencer | Illumina, San Diego, California, USA | N/A |
| Deposited Data | | |
| Raw target capture sequence data | This study | NCBI SRA PRJNA806513 |
| Raw target capture sequence data | Nikolov et al. ³ | NCBI SRA PRJNA518905 |
| Raw target capture sequence data | Hendriks et al. ⁴⁶ | NCBI SRA PRJNA678873 |
| Raw chloroplast sequence data | Walden et al. ² | ENA/GenBank PRJEB38700 |
| Multiple sequence alignments | This study | https://doi.org/10.5281/zenodo.8214354 |
| Oligonucleotides | | |
| myBaits Custom 20-40K | Arbor Biosciences, Ann Arbor, Michigan, USA | Cat#300296R |
| myBaits Angiosperms-353 | Arbor Biosciences, Ann Arbor, Michigan, USA | Cat#308196 |
| Software and Algorithms | | |
| Various scripts to map raw data, create multiple sequence alignments, gene trees, and species trees | This study | https://doi.org/10.5281/zenodo.8214354 |
| Trimmomatic v0.38 | Bolger et al. ⁸⁸ | https://github.com/usadellab/Trimmomatic |
| HybPiper v1.3.1 | Johnson et al. ⁸⁹ | https://github.com/mossmatters/HybPiper/wiki/HybPiper-Legacy-Wiki |
| BWA v0.7.16a | Li and Durbin ⁹⁰ | https://github.com/lh3/bwa |
| BWA v0.7.17 | Li and Durbin ⁹⁰ | https://github.com/lh3/bwa |
| SPAdes v3.14.1 | Bankevich et al. ⁹¹ | https://github.com/ablab/spades |
| GNU Parallel | Tange ⁹² | https://github.com/martinda/gnu-parallel |
| Python script 'filter_megatarget.py' | McLay et al. ⁹³ | See publication |
| SAMtools v1.3.1 | Danecek et al. ⁹⁴ | https://github.com/samtools/samtools |
| Picard tools | N/A | http://broadinstitute.github.io/picard/ |
| GATK3 | McKenna et al. ⁹⁵ | https://github.com/broadinstitute/gatk |
| GATK4 | Van der Auwera and O'Connor ⁹⁶ | https://github.com/broadinstitute/gatk |
| Shell script 'masker.sh' | This study | https://doi.org/10.5281/zenodo.8214354 |
| Shell script 'cpanno.py' | This study | https://doi.org/10.5281/zenodo.8214354 |
| BEDTools v2.27.1 | Quinlan and Hall ⁹⁷ | https://github.com/ark5x/bedtools2 |
| MAFFT v7.273 | Katoh and Standley ⁹⁸ | https://mafft.cbrc.jp/alignment/software/ |
| FastTree v2.1 | Price et al. ⁹⁹ | http://www.microbesonline.org/fasttree/ |
| PASTA v1.8.6 | Mirarab et al. ¹⁰⁰ | https://github.com/smirarab/pasta |
| ASTRAL-III v5.7.8 | Zhang et al. ⁴⁸ | https://github.com/smirarab/ASTRAL |
| HybPhaser v2.0 | Nauheimer et al. ⁴⁷ | https://github.com/LarsNauheimer/HybPhaser |
| SplitsTree4 v4.17.1 | Huson and Bryant ⁶⁹ | https://github.com/husonlab/splitstree4 |

(Continued on next page)

Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|--------------------------------|---|---|
| IQ-TREE2 v2.1.3 | Minh et al. ¹⁰¹ | https://github.com/iqtree/iqtree2 |
| trimAl v1.2 | Capella-Gutiérrez et al. ¹⁰² | https://github.com/inab/trimal |
| TAPER v1.0.0 | Zhang et al. ¹⁰³ | https://github.com/chaoszhang/TAPER |
| TreeShrink v1.3.9 | Mai and Mirarab ¹⁰⁴ | https://github.com/uym2/TreeShrink |
| ASTRAL-Pro v1.1.6 | Zhang et al. ⁴⁹ | https://github.com/chaoszhang/A-pro |
| R package 'ggtree' | Xu et al. ¹⁰⁵ | https://github.com/YuLab-SMU/ggtree |
| Rate4Site v3.2 | Mayrose et al. ¹⁰⁶ | N/A |
| R package 'PhyInformR v1.0' | Dornburg et al. ¹⁰⁷ | https://github.com/carolinafishes/PhyInformR |
| Perl script 'catfasta2phym.pl' | N/A | https://github.com/nylander/catfasta2phym.pl |
| IQ-TREE v1.6.12 | Nguyen et al. ¹⁰⁸ | https://github.com/Cibiv/IQ-TREE |
| IQ-TREE v2.2.0 | Minh et al. ¹⁰¹ | https://github.com/Cibiv/IQ-TREE |
| treePL v1.0 | Smith and O'Meara ¹⁰⁹ | https://github.com/blackrim/treePL |
| TreeAnnotator v2.4.7 | Bouckaert et al. ¹¹⁰ | https://beast.community/treeannotator |
| TreeAnnotator v2.6.7 | Bouckaert et al. ¹¹¹ | https://beast.community/treeannotator |

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Kasper P. Hendriks (kasper.hendriks@naturalis.nl).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Target capture data have been deposited to the NCBI Sequence Read Archive (SRA): BioProject PRJNA678873 and PRJNA806513; all data are publicly available as of the date of publication. Sample accession numbers are listed in [Table S1](#).
- All original code has been deposited to Zenodo and accessible through <https://doi.org/10.5281/zenodo.8214354>.
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

This study did not use any experimental model or study participants.

METHOD DETAILS

Taxon sampling

To reconstruct the nuclear Brassicaceae phylogeny, we aimed to include at least one species from each of the 349 currently accepted genera—preferably each genus' type species—along with one species of every non-Brassicaceae Brassicales families and all species needed for fossil calibration checks of vicariance and colonisation within lineages of Brassicaceae ([Data S1J](#)). We followed BrassiBase¹¹² for taxonomic delimitation of species, genera, and tribes, added with more recent insights from taxonomic experts. Where possible, we sampled type specimens to support future taxonomic judgements ([Table S1](#)).

We generated new nuclear sequence data for 365 samples ([Table S1](#)) and added available sequences from 38 samples from Nikolov et al.³ ([Table S2](#)). All new data were sequenced from dried herbarium specimens or silica dried tissue from 29 different herbarium collections across the world, with plants collected between 1807 and 2020 (including 35 pre-1900 and 29 samples collected between 1900 and 1950; [Methods S1B](#)). Old samples were included either because we did not have access to younger material, or to show the possibilities of our methodology with regard to natural history collections. We used the original type material of 24 species ([Table S1](#)).

New plastome data were generated from genome spiking (see below) for 237 samples ([Table S1](#)). We used additional data for 196 samples from Walden et al.,² 60 plastid genomes downloaded from GenBank and used in the same study ([Table S7](#)), and 31 samples from Nikolov et al.³ ([Table S2](#)).

Library preparation, target capture, and sequencing

Wet lab methods followed Hendriks et al.⁴⁶ Briefly, we extracted genomic DNA from 25 mg of dried leaf tissue (or less if insufficient material was available; in case no leaf tissue was available from any herbarium voucher available to us, we used branches and/or flowers) using the DNeasy PowerPlant Pro Kit (Qiagen, Hilden, Germany), following the manufacturer's protocol (but with a final elution time of 1 h). DNA extracts with visible impurities (green or brown colour; ~25% of samples) were subsequently purified using the DNeasy PowerClean Pro Cleanup Kit (Qiagen, Hilden, Germany). Genomic DNA was stored in the DNA bank of Naturalis Biodiversity Center, Leiden, the Netherlands (Table S1). Genomic libraries were generated using the NEBNext Ultra II FS kit (New England Biolabs, Ipswich, Massachusetts, USA) with sonication in an M220 Focused-ultrasonicator (Covaris, Woburn, Massachusetts, USA; only for libraries with fragment peak length > 400 bp). Indexing was performed with 384 unique combinations from IDT10 primers (Integrated DNA Technologies, Coralville, Iowa, USA), with protocol adjustments described by Hendriks et al.⁴⁶ Target sequence capture was carried out on pools of 10–30 libraries each, using the 'mixed baits' approach described by Hendriks et al.⁴⁶ This method targets putatively single-copy nuclear genes from two different bait sets in a single capture reaction: a Brassicaceae-specific set targeting 1,827 exons from 764 genes, using 40k probes³ (hereafter B764), and the now widely used Angiosperms353 v1 universal bait set targeting 353 genes, using 80k probes^{44,45} (hereafter A353; both kits available from Arbor Biosciences, Ann Arbor, Michigan, USA). To maintain the ratio of probes among the bait sets during target capture, we used a B764: A353 = 1 : 2 (v/v) mixture. To aid skimming of chloroplast gene reads during sequencing,¹¹³ we used genome spiking of each enriched library with its unenriched library at a ratio of 1 : 1 (M/M). Sequencing was performed on an Illumina NovaSeq 6000 sequencer (Illumina, San Diego, California, USA) at BaseClear, the Netherlands, producing 150-bp paired-end reads, at a targeted 100× technical coverage. Raw sequence data files were uploaded to the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) under BioProjects PRJNA806513 and PRJNA678873. Target capture and sequencing for a sample were repeated if results from sequence assembly (described below) were poor (< 500 genes recovered).

New 'mixed baits' data for 12 outgroup species were generated by the Plant and Fungal Trees of Life project at the Royal Botanic Gardens, Kew, UK, following methods described by Baker et al.¹¹⁴ A single sample was processed by the Bailey Lab, New Mexico State University, USA, and three more at Heidelberg University, Germany. To further increase sampling, we added previously published raw sequence data for 37 samples from Nikolov et al.,³ available from NCBI SRA as BioProject PRJNA518905 (target capture for B764 only; Table S2).

Sequence assembly of target capture data

Raw sequence data (from our study, as well as from other sources; see taxon sampling) were quality controlled and trimmed with Trimmomatic v0.38⁸⁸ using parameters from Baker, et al.¹¹⁴ Trimmed reads were mapped against two reference files (i.e., for B764 and A353 bait sets) using HybPiper v1.3.1⁸⁹ with BWA v0.7.16a⁹⁰ and SPAdes v3.14.1,⁹¹ using GNU Parallel⁹² to manage parallel computing of samples on the XSEDE Stampede2 HPC.¹¹⁵ We built a gene reference file for the B764 dataset from the 1,827 exon reference file of Nikolov et al.³ by concatenation of same-gene exons, resulting in a total target length of 919,712 bp. For the A353 dataset, we used the 'mega353' target file with the script 'filter_megatarget.py' to create a mustard family-specific reference⁹³ with a total target length of 294,516 bp. We identified a total of 36 genes with overlap among the two bait sets, which was expected because the two bait sets have been developed independently (Data S1K). To avoid studying the same genetic marker twice, we discarded these genes from the B764 dataset (generally the shorter targets), leaving a final nuclear dataset of 1,081 (i.e., 764 + 353 - 36) genes with a total target length of 888,392 bp.

Sequence assembly of plastome

We used off-target reads from genome spiking (samples with new raw data) and genome skimming (raw data from previous studies listed above) to reconstruct 60 plastid genes for 237 new samples and 31 samples from Nikolov et al.³ New data were integrated into an already existing plastid dataset containing 231 Brassicaceae species.²

Trimmed sequencing reads were mapped using BWA v0.7.17 using option 'BWA-MEM'¹¹⁶ and the *Arabidopsis thaliana* plastid genome (NCBI GenBank accession number NC_000932) as reference. Prior to mapping, the second copy of the inverted repeat region was removed, as identical regions lead to secondary alignments which are omitted by tools used in subsequent analysis. SAMtools v1.3.1⁹⁴ was used to enhance mapping quality and to sort and index the .bam files. Duplicates were removed using Picard tools (<http://broadinstitute.github.io/picard/>). Variant calling was performed using the GATK4⁹⁶ function 'HaplotypeCaller' setting ploidy to 1 and pcr-indel-model to none. The GATK3⁹⁵ function 'FastAlternateReferenceMaker' was used to generate sequences including the detected SNPs and indels. Regions of low coverage (< 5) and low mapping quality (< 30) were detected using GATK3 function 'CallableLoci'. After having adjusted the positions of the regions to be masked using the inhouse script 'masker.sh' (Markus Kiefer, Heidelberg University; see Key Resources Table), BEDTools⁹⁷ function 'maskfasta' was used to mask regions of bad mapping quality (< 30) and low coverage (< 5). The annotation of genes was transferred by alignment to above mentioned *A. thaliana* plastid reference using the inhouse script 'cpanno.py' (Markus Kiefer, Heidelberg University; see Key Resources Table). After removal of gap columns, the final data matrix had a total target length of 29,120 bp and was 96.2% complete.

Taxonomic verification

We performed 'taxonomic verification' on a preliminary species phylogeny. To reconstruct this phylogeny, multiple sequence alignments were created for each gene using MAFFT v7.273,⁹⁸ with a quick gene tree inference using FastTree v2.1,⁹⁹ both within the

pipeline PASTA v1.8.6.¹⁰⁰ Unfiltered gene trees were used as input to ASTRAL-III v5.7.8.⁴⁸ Data from samples marked as possible or likely errors (found in highly unlikely positions in the preliminary species tree) were removed, followed by either repetition of library preparations or resampling of the species. This routine was repeated once more, such that the phylogenetic position of each sample was verified by taxonomic experts. Subsequently, any data resulting from multiple DNA extractions and/or library preparations from the same voucher were merged after trimming and again mapped following the same routine.

Allelic variation and paralog detection

We used HybPhaser v2.0,⁴⁷ an extension to HybPiper, to assess allelic variation and to detect possible paralogs in our nuclear dataset. In short, HybPhaser performs a re-mapping of raw sequence data, using the contig of each sample (created by HybPiper) as a new reference. Whereas HybPiper by default constructs the most likely allele for a—supposedly single-copy—gene based on the relative nucleotide frequency of each heterozygous site, HybPhaser instead takes SNP variation into account using nucleotide ambiguity codes, and uses these to quantify divergence between gene variants to detect paralogy and hybridisation. Single genes with high SNP count are considered likely paralogs, while samples with high SNP count across all genes are considered likely hybrids or polyploids.⁴⁷ Putative paralogs were removed from the dataset. However, since there is no single criterion to define a paralog, we used five ‘routines’ to do this (Table 1). In the ‘inclusive’ routine (1,018 genes, 332 genera, 375 species, after running a de-noising loop; see below), we retained as much data as possible (including all samples, and thus all genera for which we had any data) and only discarded poorly recovered genes (gene recovered for < 10% of the samples and/or proportion of gene target length recovered < 10% on average across all samples). We discarded putative paralogs in the ‘strict’ routine (1,013 genes, 317 genera, 356 species) by removing all ‘outlier’ genes, defined as loci that have more than 1.5*IQR (interquartile range) above the 3rd quartile of mean SNPs. In a ‘superstrict’ routine (297 genes, 317 genera, 356 species), we removed all genes with a mean proportion of SNPs across the dataset of > 0.02. After noticing large differences in mean SNP proportions among tribes within the mustard family (Methods S1C), we added the ‘superstrict by tribe’ routine (mean 1,031 genes, with gene selection varying by tribe, 317 genera, 356 species) in which we assessed and removed mean SNP proportions by tribe, leading to a very sparse sample-gene matrix (Methods S1D). Finally, we aimed to improve phylogenetic backbone support further by using a ‘superstrict excluding hybrids’ routine (303 genes, 124 genera, 138 species). This dataset was the same as in the ‘superstrict’ routine, but all samples belonging to rogue taxa and/or having a locus heterozygosity and/or allelic divergence (see next) in the upper 50% of detected values from all samples were removed. This led to a highly reduced (in terms of both genes and samples/species/genera/tribes) dataset, but with the anticipated advantage of having removed as much noise from the dataset as possible, while at the same time including enough representatives from all main lineages.

We used HybPhaser to detect possible hybrids by calculating each sample’s allele divergence (AD; percentage of SNPs across all genes) and locus heterozygosity (LH; percentage of genes with SNPs), two metrics that are useful in the detection of hybrids.⁴⁷ Because hybrids are expected to inherit multiple alleles from their different parent species, they are expected to show relatively high levels of LH and an AD that corresponds to the divergence of the parental lineages. Very high values for AD are expected in lineages with multiple polyploidizations.⁴⁷ With time, polyploid lineages are expected to lose duplicated genes leading to a decrease in LH. Therefore, high AD combined with intermediate LH can indicate that samples are more ancient polyploids. While there is no universal circumscription of what values correspond to hybrids or other types of polyploids, these values can give a good indication on the history of hybridisation in samples. Here we broadly distinguish four classes: hybrid (high LH, medium AD), highly polyploid (high LH and high AD), old polyploid (medium LH, medium AD), and old and highly polyploid (medium LH, high AD).

Nuclear phylogenomics

We applied four different phylogenomic approaches to analyse our nuclear dataset. We used default settings and parameters for all tools, unless specified.

First, we applied a network approach to visualise possible evolutionary reticulations, inferring a splits graph (based on uncorrected p-distances) with SplitsTree4 v4.17.1.⁶⁹ We used a nuclear supermatrix from the 297 gene alignments from the ‘superstrict’ routine as input.

Second, we used a ML supermatrix approach with IQ-TREE2 v2.1.3¹⁰¹ with again the nuclear supermatrix approach with the 297 genes from the ‘superstrict’ routine (with outgroup sample S1321, *Synsepalum afzellei*), and separately with a supermatrix of the 303 genes from the ‘superstrict excluding hybrids’ routine (with outgroup sample MYZV, *Tropaeolum peregrinum*). We used 1,000 ultra-fast bootstraps¹¹⁷ (saving bootstrap replicates) with a GTR+F+R model. In this first analysis we specifically kept all outgroup samples needed for fossil calibration, even if these would have been removed based on our sampling routine criteria. Contrary to our coalescent-based approach (see below), these analyses generated phylogenies in which branch lengths were representative of evolutionary change (number of mutations), which was needed for the subsequent divergence time estimation. As an input for fossil calibration (see below) we again used a nuclear supermatrix approach with the 297 genes from the ‘superstrict’ routine. We repeated the IQ-TREE2 analysis with the resultant species tree and all 297 gene trees (see below) to calculate gene (gCF) and site concordance factors (sCF; parameter –scf 1,000) for all nodes.⁶⁷

Third, we applied a coalescent-based approach with ASTRAL-III.⁴⁸ As input, we took the consensus sequences from the four paralog detection routines in HybPhaser (see above) to infer gene trees that served as input for a coalescent-based analysis. For each of the routines, we started by running a de-noising loop: sequences were aligned using MAFFT v7.273⁹⁸ and trimmed using trimAl v1.2¹⁰² with parameters resoverlap 0.75, seqoverlap 0.90, and gt 0.90. Any remaining likely sequencing errors were masked using

TAPER v1.0.0¹⁰³ with default parameters. Gene trees were inferred using IQ-TREE2 v2.1.3¹⁰¹ inferring branch support using ultrafast bootstrapping,¹¹⁷ with other parameters following Baker, et al.¹¹⁴ We used TreeShrink v1.3.9¹⁰⁴ with default parameters on the complete set of gene trees to detect and remove outlier branches and update gene trees and alignments. Discordance among gene trees was scored using all gene trees associated with each routine and calculated using normalised quartet scores for the main topology, along with first and second alternatives.

Fourth, we applied another coalescent-based approach with ASTRAL-Pro v1.1.6.⁴⁹ Contrary to ASTRAL-III, this version allows the input of multiple alleles (including possible paralogs) for each individual, acknowledging that this may actually be informative in species tree inference (e.g., no a priori choice of a definitive homologous gene copy needs to be made). Gene trees were now collected from mapping done by HybPiper using the script ‘paralog_investigator.py’, which saves all possible alleles.⁸⁹ Genes were again aligned with MAFFT v7.273 and trimmed using trimAl v1.2 with parameters resoverlap 0.75, seqoverlap 0.90, and gt 0.90, with subsequent gene tree inference with IQ-TREE2 v2.1.3. All phylogenetic trees from the approaches 2–4 were plotted using the R package ggtree.¹⁰⁵

Phylogenetic informativeness

To study any differences in support from different nuclear genes in inferring the nuclear species tree, we calculated Townsend’s phylogenetic informativeness⁵⁰ for all genes included in the ‘strict’ routine (Figure S1). First, we calculated relative evolutionary rates for all sites in each gene’s multiple sequence alignment, constrained on the ML supermatrix approach species tree, using Rate4Site v3.2.¹⁰⁶ Second, we used the R package PhyInformR v1.0¹⁰⁷ to calculate phylogenetic informativeness profiles for each gene, making a distinction between genes obtained from either the B764 and A353 bait sets, and genes with a mean SNP proportion of ≤ 0.02 and > 0.02 (i.e., the threshold applied for paralog detection in HybPhaser).

Plastome phylogenetics

To generate a plastome-based phylogeny, coding sequences and sequences encoding tRNAs and rRNAs were extracted using BEDTools v2.27.1 function ‘getfasta’⁹⁷. We used a reduced gene set of 60 loci as previously used by Walden et al.² New sequences were aligned, together with the corresponding sequences from Walden et al.,² using MAFFT v7.45.3.⁹⁸ In a last step, gap columns were deleted and alignments were concatenated using the script ‘catfasta2phym.pl’ (<https://github.com/nylander/catfasta2phym>).

Phylogenetic reconstruction was performed using IQ-TREE v1.6.12¹⁰⁸ with partition information from the alignment, defining the outgroup, and running 1,000 ultrafast bootstrap replicates.¹¹⁷ A second phylogenetic reconstruction was performed using IQ-TREE v2.2.0¹⁰¹ to calculate site concordance factors (sCF) as a measure of support for the splits in the tree,⁶⁷ with the former species tree topology and gene alignment as input. Note that in the plastome phylogeny, gene concordance factors (gCF) cannot be calculated as in the nuclear phylogeny, because the 60 plastome genes studied were considered to be a single heritable unit (supermatrix approach), and consequently separate gene trees (needed to calculate gCF) were not inferred.

Nuclear versus plastome BrassiToL species tree incongruences were visualised using a cophylogeny plot created in R package phytools.¹¹⁸ Incongruences within tribes were quantified using the classical Robinson-Foulds metric¹¹⁹ and the generalised Robinson-Foulds metric following Smith¹²⁰ using R package TreeDist.¹²⁰ Species trees were first pruned to include only species present in both trees, and any duplicates were removed (after random selection per species).

Fossil calibration

For both the nuclear and the plastome dataset, we estimated divergence times within the family using the penalized likelihood approach¹²¹ as implemented in treePL v1.0¹⁰⁹ and the Turonian fossil *Dressiantha bicarpellata*, estimated at 93.6–89.3 Ma⁵¹ (which we took as maximum and minimum ages, respectively), as a single calibration point at the stem node of order Brassicales (cf. Couvreur et al.²⁰). For the nuclear dataset, we used the topology from the ML supermatrix approach as input species tree, and reran IQ-TREE using the gene alignments from the 20 most clock-like genes only to infer relative branch lengths, acknowledging that inclusion of too many genes can easily result in an artificial pushback in time of internal nodes. To do so, we first calculated the clock-likeness of all genes following Vankan et al.,¹²² who defined clock-likeness as the coefficient of variation of all root-to-tip distances in the gene tree. When running the priming analysis in treePL, the value for ‘opt’ was set to 2, and ‘optad’ set to 1. The ‘moredetail’ and ‘moredetailad’ options were in effect and ‘optcvad’ was set to 1. Cross validation analysis indicated 10 as the best smoothing value. We assessed node age estimates by repeating the treePL (using the above optimised settings) analysis for 1,000 bootstrap trees generated with IQ-TREE, this time fixing the topology of the species tree (but not branch lengths) and summarising with TreeAnnotator v2.4.7¹¹⁰ to obtain 95% HPD confidence intervals.

For the plastid dataset, when running the priming analysis and later adjustments, the values for ‘opt’ and ‘optad’ were both set to 3. The ‘moredetail’ and ‘moredetailad’ options were in effect and ‘optcvad’ was set to 4. Cross validation analysis indicated 0.00001 as best smoothing value. Again, we assessed node age estimates by repeating the treePL analysis (using the above optimised settings) for the 1,000 bootstrap replicates generated in IQ-TREE. Calibrated gene trees were summarised using TreeAnnotator v2.6.7¹¹¹ to obtain 95% HPD confidence intervals.

We performed multi-evidence validation of our new results against four other fossils and five biogeographical dating events as suggested by Franzke et al.⁵² by comparing expected and recovered node ages (Data S1J).

QUANTIFICATION AND STATISTICAL ANALYSIS

Support for nodes in phylogenetic trees was calculated using multiple methods (LPP, bootstrap, quartet score, gCF, and sCF), as detailed under [method details](#).