
































# Complementing model species with model clades

Makenzie E. Mabry <sup>1</sup>, R. Shawn Abrahams <sup>2,3</sup>, Ihsan A. Al-Shehbaz <sup>4</sup>, William J. Baker <sup>5</sup>,  
Simon Barak <sup>6</sup>, Michael S. Barker <sup>7</sup>, Russell L. Barrett <sup>8</sup>, Aleksandra Beric <sup>9,10</sup>,  
Samik Bhattacharya <sup>11</sup>, Sarah B. Carey <sup>12</sup>, Gavin C. Conant <sup>13</sup>, John G. Conran <sup>14</sup>,  
Maheshi Dassanayake <sup>15</sup>, Patrick P. Edger <sup>16</sup>, Jocelyn C. Hall <sup>17</sup>, Yue Hao <sup>18</sup>,  
Kasper P. Hendriks <sup>11,19</sup>, Julian M. Hibberd <sup>20</sup>, Graham J. King <sup>21</sup>, Daniel J. Kliebenstein <sup>22</sup>,  
Marcus A. Koch <sup>23</sup>, Ilia J. Leitch <sup>5</sup>, Frederic Lens <sup>19,24</sup>, Martin A. Lysak <sup>25</sup>, Alex C. McAlvay <sup>26</sup>,  
Michael T.W. McKibben <sup>7</sup>, Francesco Mercati <sup>27</sup>, Richard C. Moore <sup>28</sup>, Klaus Mummenhoff <sup>11</sup>,  
Daniel J. Murphy <sup>29</sup>, Lachezar A. Nikolov <sup>30</sup>, Michael Piasis <sup>31</sup>, Eric H. Roalson <sup>32</sup>,  
M. Eric Schranz <sup>33</sup>, Shawn K. Thomas <sup>34,35</sup>, Qingyi Yu <sup>36</sup>, Alan Yocca <sup>12</sup>, J. Chris Pires <sup>37</sup>  
and Alex E. Harkess <sup>12,\*</sup>

- 1 Florida Museum of Natural History, University of Florida, Gainesville, FL 32611, USA
- 2 Department of Ecology and Evolutionary Biology, Yale University, New Haven, CT 06520, USA
- 3 Department of Biochemistry, Purdue University, West Lafayette, IN 47906, USA
- 4 Missouri Botanical Garden, Shaw Boulevard, St. Louis, MO 63110, USA
- 5 Royal Botanic Gardens, Kew, Richmond, Surrey TW9 3AE, UK
- 6 Ben-Gurion University of the Negev, French Associates Institute for Agriculture and Biotechnology of Drylands, Jacob Blaustein Institutes for Desert Research, Midreshet Ben-Gurion, 8499000, Israel
- 7 Department of Ecology and Evolutionary Biology, University of Arizona, Tucson, AZ 85721, USA
- 8 National Herbarium of New South Wales, Australian Botanic Garden, Locked Bag 6002, Mount Annan, NSW 2567, Australia
- 9 Department of Psychiatry, Washington University in Saint Louis School of Medicine, St. Louis, MO 63110, USA
- 10 NeuroGenomics and Informatics Center, Washington University in Saint Louis School of Medicine, St. Louis, MO 63108, USA
- 11 Department of Biology, Botany, University of Osnabrück, D-49076 Osnabrück, Germany
- 12 HudsonAlpha Institute for Biotechnology, Huntsville, AL 35806, USA
- 13 Department of Biological Sciences, Bioinformatics Research Center, Program in Genetics, North Carolina State University, Raleigh, NC 27695, USA
- 14 ACEBB and SGC, School of Biological Sciences, The University of Adelaide, Adelaide, SA 5005, Australia
- 15 Department of Biological Sciences, Louisiana State University, Baton Rouge, LA 70803, USA
- 16 Department of Horticulture, Michigan State University, East Lansing, MI 48864, USA
- 17 Department of Biological Sciences, University of Alberta, Edmonton, Alberta T6G 2E9, Canada
- 18 Cancer and Cell Biology Division, Translational Genomics Research Institute, Phoenix, AZ 85004, USA
- 19 Functional Traits, Naturalis Biodiversity Center, PO Box 9517, Leiden 2300 RA, the Netherlands
- 20 Department of Plant Sciences, University of Cambridge, Cambridge, CB2 1TN, UK
- 21 Southern Cross Plant Science, Southern Cross University, Lismore, NSW 2480, Australia
- 22 Department of Plant Sciences, UC Davis, Davis, CA 95616, USA
- 23 Centre for Organismal Studies (COS), Heidelberg University, 69120 Heidelberg, Germany
- 24 Institute of Biology Leiden, Plant Sciences, Leiden University, 2333 BE Leiden, the Netherlands
- 25 CEITEC, and NCBR, Faculty of Science, Masaryk University, 625 00 Brno, Czech Republic
- 26 Institute of Economic Botany, New York Botanical Garden, The Bronx, NY 10458, USA
- 27 National Research Council (CNR), Institute of Biosciences and Bioresource (IBBR), Palermo 90129, Italy
- 28 Biology Department, Miami University, Oxford, OH 45056, USA
- 29 Royal Botanic Gardens Victoria, Melbourne, VIC 3004, Australia

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- 30 Department of Biology, Indiana University, Bloomington, IN 47405, USA  
31 Division of Plant Sciences, University of Missouri, Columbia, MO 65211, USA  
32 School of Biological Sciences, Washington State University, Pullman, WA 99164-4236, USA  
33 Biosystematics Group, Wageningen University, 6708 PB Wageningen, the Netherlands  
34 Division of Biological Sciences, University of Missouri, Columbia, MO 65211, USA  
35 Bioinformatics and Analytics Core, University of Missouri, Columbia, MO 65211, USA  
36 Daniel K. Inouye U.S. Pacific Basin Agricultural Research Center, Agricultural Research Service, United States Department of Agriculture, Hilo, HI 96720, USA  
37 Department of Soil and Crop Sciences, Colorado State University, Fort Collins, CO 80523-1170, USA

\*Author for correspondence: aharkess@hudsonalpha.org

## Abstract

Model species continue to underpin groundbreaking plant science research. At the same time, the phylogenetic resolution of the land plant tree of life continues to improve. The intersection of these 2 research paths creates a unique opportunity to further extend the usefulness of model species across larger taxonomic groups. Here we promote the utility of the *Arabidopsis thaliana* model species, especially the ability to connect its genetic and functional resources, to species across the entire Brassicales order. We focus on the utility of using genomics and phylogenomics to bridge the evolution and diversification of several traits across the Brassicales to the resources in *Arabidopsis*, thereby extending scope from a model species by establishing a “model clade.” These Brassicales-wide traits are discussed in the context of both the model species *Arabidopsis* and the family Brassicaceae. We promote the utility of such a “model clade” and make suggestions for building global networks to support future studies in the model order Brassicales.

## Introduction

Model species are essential in plant biology. For example, Barbara McClintock carefully tracked corn (*Zea mays*) kernel variation to characterize transposable elements and chromosome segregation (McClintock 1941, 1950). The genome of *Arabidopsis thaliana* (The *Arabidopsis* Genome Initiative 2000) and foundational mutant and germplasm collections like the *Arabidopsis* T-DNA Insertional Mutant Collection (O’Malley et al. 2015) have driven our understanding of gene function in all plants. However, the plant kingdom is vast, comprising nearly 500,000 species and spanning a billion years of evolutionary time (Nic Lughadha et al. 2016). Collaborations among systematists, herbaria, and botanical gardens continue to drive advances in building the plant tree of life (Johnson et al. 2019; One Thousand Plant Transcriptomes Initiative 2019), in part due to rapid advances and the decreasing cost of genome and transcriptome sequencing technologies. These technological advancements enable us to build resources that complement and elevate the usefulness of a model species, encompassing larger taxonomic groups to now produce “model clades” of plants. As species relationships become clearer, the plant biology community is in an ideal position to leverage comparative genomics across these clades to unravel the evolution and function of genes, regulatory networks, repetitive elements, and noncoding DNA. Comparative biologists and phylogeneticists have long been interdisciplinarily working on these problems, for example, to examine floral organ evolution across the order Ranunculales (Damerval and Becker 2017) and the grasses (Schragger-Lavelle et al. 2017). The development of accessible comparative genome and gene family

evolution platforms like Phytozome (Goodstein et al. 2012), CoGE (Lyons 2008), Ensembl Plants (Yates et al. 2022), and GENESPACE (Lovell et al. 2018, 2022) allow the placement of stand-alone genome assemblies in a larger evolutionary and phylogenetic context, creating an opportune time for plant biologists to identify clades with diverse phenotypes that can be anchored to a closely related model species.

The order Brassicales exhibits extensive diversity of species, traits, and environmental adaptations across a broad geographical distribution and includes several crops like canola, caper, broccoli, kale, papaya, rapeseed, and saltwort. Within the Brassicales, one can find recurrent evolution and diversification of complex traits involving genomics (e.g. hybridization, polyploidy, repeat sequence turnover, speciation, and reproductive systems), physiology (e.g. extremophytism, photosynthesis modes), morphology (e.g. woodiness, tuberization), metabolomics (e.g. oil production, secondary metabolite diversification), and environment (e.g. wet and hyperarid habitats). To understand the genetic underpinnings of these traits, the predicted functions of orthologous genes in any understudied Brassicales species can be anchored to the massive mutant collections in the Brassicaceae “model species” *A. thaliana*. In this review, we highlight the utility of all Brassicales for benefiting from comparative genomics to understand trait variation and evolution in the context of producing a “model clade” of plants that builds out from a “model species.” We frame each major section with 3 key points. First, we describe what is known about a given trait from studying the family Brassicaceae, often derived from research using *A. thaliana*. Second, we describe what is known

about the diversity or variation of a given trait across the order Brassicales. Third, we discuss how leveraging comparative genomics across the order will lead to a better understanding of gene, genome, and trait evolution.

## Systematics and diversification of the Brassicales

### Systematics across the Brassicaceae

The Brassicales comprise ca. 4,700 species in 18 families (Table 1; Magallon et al. 1999; Swanepoel et al. 2020), with the Brassicaceae, by far the largest family, comprising ca. 4,140 species in 349 genera (Koch et al. 2018; Walden et al. 2020a; German et al. 2023; Hendriks et al. 2023; Table 1). Brassicaceae systematics has long been complicated by high levels of parallel and convergent evolution of morphological characters, likely reflecting recent and rapid radiations (Huang et al. 2016; Guo et al. 2017; Walden et al. 2020a; Zuo et al. 2022). Nearly all species are assigned to 1 of 58 monophyletic tribes (German et al. 2023; Hendriks et al. 2023), which in turn are assigned to 2 subfamilies: the small monogeneric Aethionemoideae and the large Brassicoideae (98.6% of species; German et al. 2023). Additionally, the 5 main lineages within Brassicoideae recently received supertribe status (German et al. 2023). While assignment of tribes to supertribes is more or less consistent between reconstruction methods, relationships among supertribes vary. For instance, in plastid phylogenies, the Camelinoideae (formerly lineage I) consistently diverges first (Fig. 1; Huang et al. 2016; Guo et al. 2017; Mabry et al. 2020; Walden et al. 2020a; Hendriks et al. 2023), whereas nuclear phylogenies show the Hesperoideae (formerly lineage III) diverging first (Fig. 1; Huang et al. 2016; Nikolov et al. 2019; Mabry et al. 2020; Hendriks et al. 2023). With the availability of whole genome sequences from all main lineages and advances in synteny and coalescent-based methods for phylogenetic inference, additional evolutionary hypotheses have been proposed recently (Walden et al. 2020b; Zou et al. 2020). For example, the similarity of the *Arabidopsis thaliana* genome structure with that of *Aethionema* (Aethionemoideae) may indicate an earlier diverging position of the lineage leading to Arabidoideae (Walden et al. 2020b) and potentially Alyseae (both supertribe Araboideae, formerly lineage IV; Hendriks et al. 2023). Such contrasting gene and species tree topologies suggest complex evolutionary histories and lead to natural complications in achieving a stable classification based on the principle of monophyly at many taxonomic levels (Forsythe et al. 2020).

### Systematics across the Brassicales

The morphological, biochemical, and diverse genomic composition across the Brassicales permits comparative investigation of evolutionary novelties and macroevolutionary patterns (Fig. 2). However, these approaches rely on first constructing a well-supported species tree. Most of the families across the

**Table 1.** The distribution of families within the Brassicales, along with the noted genera and species numbers<sup>a</sup>

Family	Genera/Species	Geographical Distribution
Akaniaceae	2/2	<i>Akania</i> – Australia <i>Bretschneidera</i> – Asia
Bataceae	1/2	Tropical America, Austral-Asia
Brassicaceae	349/4,140	Cosmopolitan (mostly temperate)
Capparaceae	16/480	Cosmopolitan (mostly tropical)
Caricaceae	6/34	Tropical Africa and America (mostly New World)
Cleomaceae	26+/270	Cosmopolitan
Emblingiaceae	1/1	Southwest Australia
Gyrostemonaceae	5/18+	Australia
Koeberliniaceae	1/2	North, Central, and South America
Limnanthaceae	2/8	North America
Moringaceae	1/12	Africa, Madagascar, and Asia
Pentadiplandraceae	1/1	Western Africa
Resedaceae	8/96	North Africa, Eurasia, North America
Salvadoraceae	3/11	Africa to Southeast Asia
Setchellanthaceae	1/1	Mexico, North America
Tiganophytaceae	1/1	Namibia, Africa
Tovariaceae	1/2	Tropical America
Tropaeolaceae	1/105	North and South America

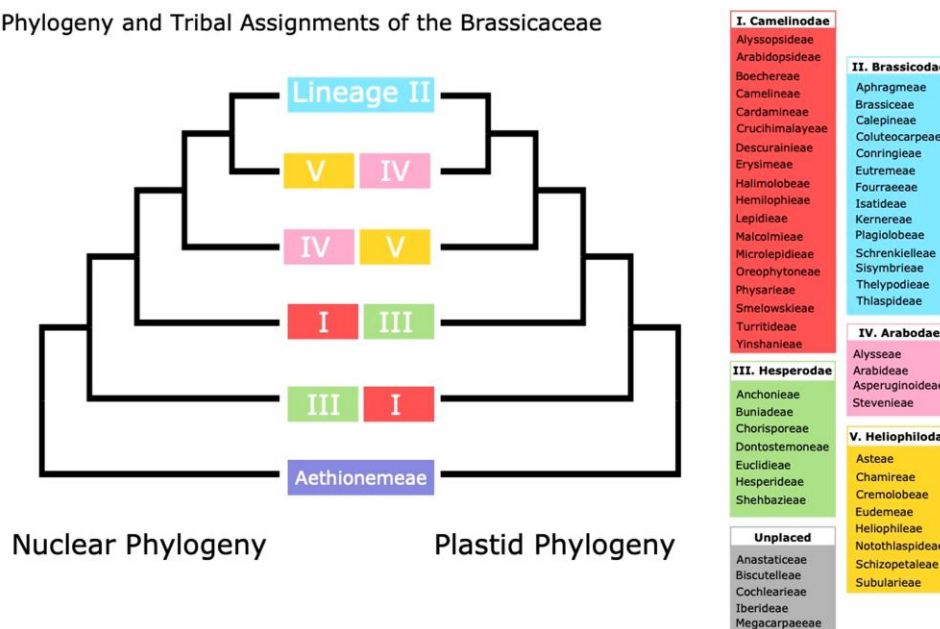
<sup>a</sup>Genera and species numbers from the Angiosperm Phylogeny Website (Stevens 2001). The recently proposed family Borthwickiaceae (found in China and Myanmar) is included within Resedaceae.

Brassicales, including Borthwickiaceae (Zhao et al. 2015) and the recently described Tiganophytaceae (Swanepoel et al. 2020), are only minimally sampled at the species level. Relationships across the order have been established with phylotranscriptomics (Edger et al. 2015, 2018a; Mabry et al. 2020), target sequence capture (Baker et al. 2022), and more traditional molecular systematic approaches based on fewer genes and larger taxonomic sampling (Rodman et al. 1993, 1996; Cardinal-McTeague et al. 2016). Many relationships among the Brassicales are well supported (Hall et al. 2002, 2004; Ronse De Craene and Haston 2006; Hall 2008; Edger et al. 2015, 2018a; Cardinal-McTeague et al. 2016; Mabry et al. 2020); however, some key nodes remain unresolved, which may be the consequence of rapid diversification and/or introgression (Edger et al. 2018a). Due to a large number of extant Brassicales species, building a high-resolution species tree requires both a unified research community effort as well as creative approaches to integrating multiple types of data.

### The future of Brassicales systematics research

A complete tree of all Brassicales species, with multiple representative sampling for each species, is desirable and tractable, especially given the growing popularity of reduced representation methods such as targeted sequence capture (Dodsworth et al. 2019; Baker et al. 2021). Targeted sequence capture can be applied very effectively even to degraded DNA obtained from herbarium specimens (Brewer et al. 2019), which vastly increases opportunities for extensive

## Phylogeny and Tribal Assignments of the Brassicaceae



**Figure 1.** Phylogeny indicating the lineage relationships of the Brassicaceae using nuclear and plastid data. Placed tribes are noted with corresponding color and numbering. Summary, classification, and relationships based on Hendriks et al. (2023).

species sampling. These methods have proven useful for understanding species relationships at the order level across angiosperms (Antonelli et al. 2021; Lee et al. 2021; Maurin et al. 2021; Thomas et al. 2021; Zuntini et al. 2021), and efforts are already underway to build comprehensive phylogenomic datasets across the Brassicales by employing a universal sequence capture toolkit (Johnson et al. 2019; Baker et al. 2022). The combination of universal and lineage-specific toolkits (Nikolov et al. 2019; Eserman et al. 2021; Hendriks et al. 2021) offers great promise for a complete species phylogeny of the order. Further, a Department of Energy Joint Genome Institute Community Sequencing Project (<https://jgi.doe.gov/csp-2021-brassicales-genome-initiative/>) is currently funded to assemble and annotate chromosome-scale assemblies for at least 26 species, spanning every family in the order. The future for the phylogenomic community studying the Brassicales likely resides in a dual approach, integrating reduced representation genomic data with high-quality, chromosome-scale genomes and annotations.

## Polyploidy and Diploidization Dynamics

### Polyploidy across the Brassicaceae

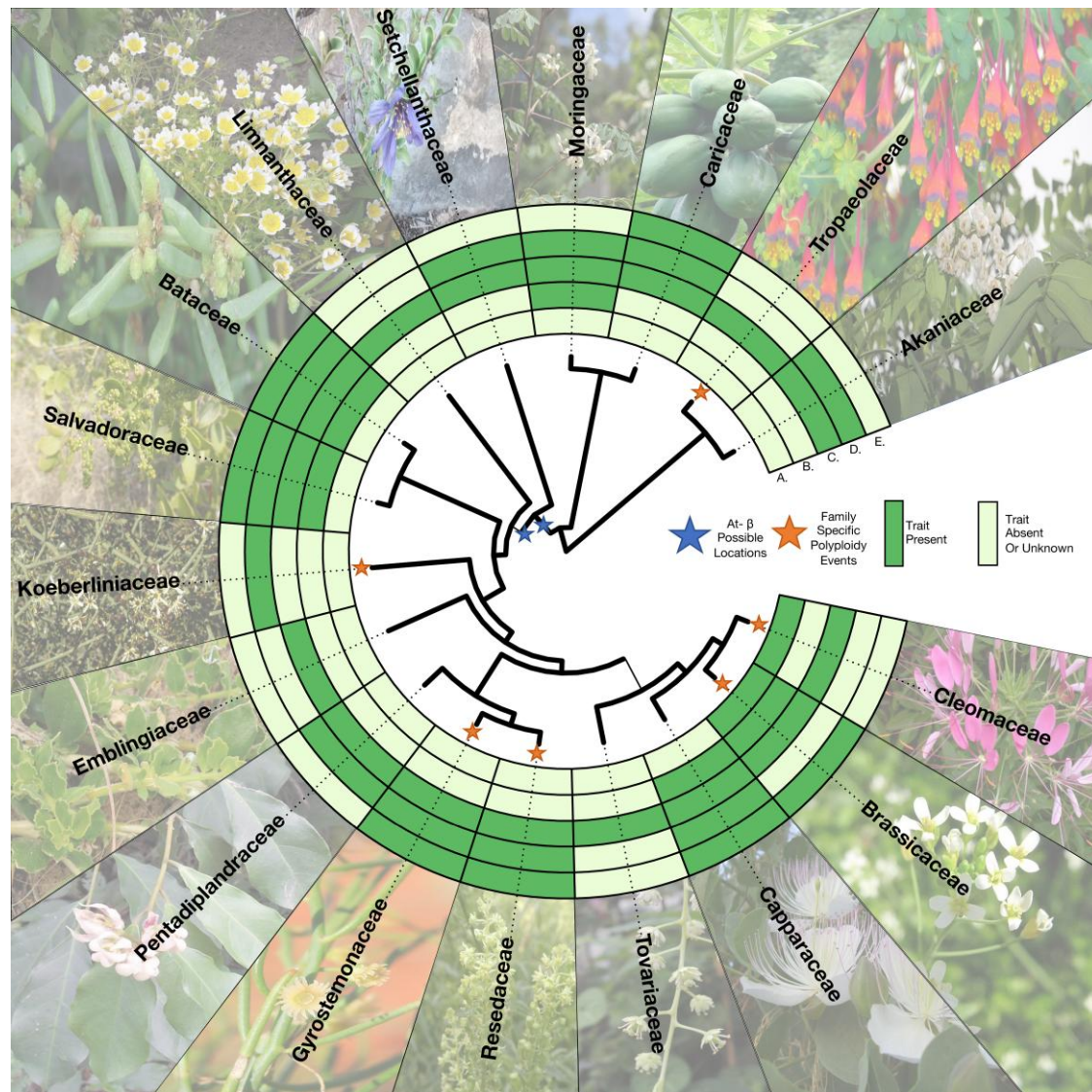
Whole-genome duplications (WGDs) are prevalent throughout plant evolution and are often associated with the origin of novel traits and shifts in diversification (Schranz et al. 2012; Van de Peer et al. 2021). Early studies suggested that *Brassica* species (Brassicaceae) were hexaploids relative to *A. thaliana* (Arabidopsiadeae) and that their genomes experienced chromosomal rearrangements during their return to the diploid state (Lagercrantz 1998; Babula et al. 2003). These results were confirmed by later studies and the sequencing of

*Brassica* genomes (Lysak et al. 2005, 2007; Parkin et al. 2005; Schranz et al. 2006; *Brassica rapa* Genome Sequencing Project Consortium 2011; Fig. 3). This hexaploidy or whole-genome triplication (WGT) occurred through 2 successive hybridizations creating first an allotetraploid and eventually an allohexaploid (Cheng et al. 2012; Tang et al. 2012). However, the timing and placement of the hexaploidy on a resolved phylogeny remains elusive (Lysak et al. 2007; *Brassica rapa* Genome Sequencing Project Consortium 2011; Arias and Pires 2012; Cardinal-McTeague et al. 2016; Huang et al. 2016). More than 13 other independent, genus- or tribe-specific, polyploidy events have been identified across the Brassicaceae (Mandáková et al. 2017; Huang et al. 2020; Walden et al. 2020b; Hendriks et al. 2023). A study of nearly one-half of all Brassicaceae species found that 50% of these are recent polyploids (i.e. still recognized as polyploids based on cytological data) with at least 1 hidden state (characters that are not observed for the purpose of the model but influence the diversification rate (Beaulieu and O'Meara 2016), which impacts the net diversification of diploid and polyploid species (Román-Palacios et al. 2020). Another study by Walden et al. (2020a) suggests these unassessed traits from Román-Palacios et al. (2020) may be associated with morphological diversity.

### Polyploidy across the Brassicales

Multiple studies have found ancient WGDs spread more broadly across the Brassicales (e.g. At- $\beta$  event; Fig. 2; Barker et al. 2009; Edger et al. 2015, 2018a; One Thousand Plant Transcriptomes Initiative 2019; Mabry et al. 2020). Many families have also experienced family-specific WGDs, including the Brassicaceae (At- $\alpha$ ; Barker et al. 2009; Haudry et al. 2013;





**Figure 2.** Phylogeny of 17 families in the Brassicales and character trait matrix. The presence of a given trait is marked by a filled dark-green square. Traits that are unknown or absent are in light green. Possible locations of At- $\beta$  are marked in blue stars, with family-specific whole-genome duplications in orange. Traits displayed are: A) C4, B) extremophytism, C) glucosinolates, D) ancestral woodiness, and E) dioecy. Additional proposed families, which are not included here, still need to be analyzed using multiple single-copy nuclear genes (e.g. Tiganophytaceae and Borthwickiaceae). Areas of the phylogeny with no or low support are indicated by thin branch lines. Topology and support based on [Edger et al. \(2018a\)](#).

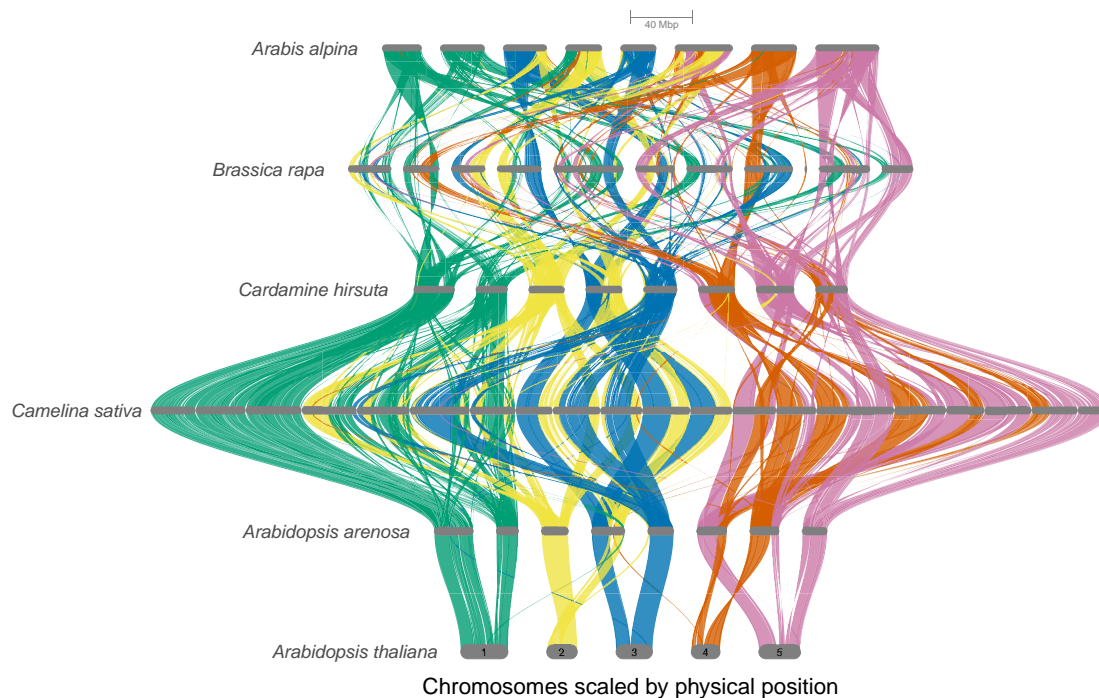
[Edger et al. 2015, 2018a](#); [One Thousand Plant Transcriptomes Initiative 2019](#); [Mabry et al. 2020](#)), Cleomaceae (Cs- $\alpha$  or Th- $\alpha$ ; [Schranz and Mitchell-Olds 2006](#); [Barker et al. 2009](#); [Bayat et al. 2018](#); [Mabry et al. 2020](#); [Hoang et al. 2023](#)), and Resedaceae (Rs- $\alpha$ ; [Lysak 2018](#); [One Thousand Plant Transcriptomes Initiative 2019](#); [Mabry et al. 2020](#)). Family-specific WGDs have also been identified in Gyrostemonaceae, Koeberliniaceae, and Tropaeolaceae ([One Thousand Plant Transcriptomes Initiative 2019](#)).

### The future of Brassicales polyploidy research

Understanding polyploidy and diploidization in the Brassicales is highly relevant and impactful for several reasons. Firstly, the Brassicales are ideal for studying these processes due to their

relatively small genomes ( $N = 342$  taxa, mean = 761.0 mb/1C, median = 537.0 mb/1C, mode = 392.0 mb/1C; based on data from the Plant DNA C-values database release 7.1 together with additional data not yet incorporated; [Pellicer and Leitch \(2020\)](#) and [Leitch et al., release 7.1, April 2019](#)). We can also produce high-quality genomes that allow for fine-scale comparisons ([Kreiner et al. 2017](#)), making them well-suited for in-depth genomic analysis. This relevance stems from the potential insights that can be gained into broader biological phenomena.

The impact of this research extends to various areas. Firstly, resolving the current struggles in understanding gene loss and retention after polyploidy in Brassicales can provide critical insights into the evolution of complex genomes. While



**Figure 3.** Whole-genome synteny of chromosome-scale Brassicaceae genomes. The synteny plot was generated using GENESPACE v0.9.1 (Lovell et al. 2022) with “diamondMode” and “orthofinderMode” set to fast, using *A. thaliana* (Arabidopsidae; Lamesch et al. 2012), *A. arenosa* (Arabidopsidae; Barragan et al. 2021), *B. rapa* (Brassicaceae; v1.3; downloaded from Phytozome), *Camelina sativa* (Camelineae; Kagale et al. 2014), *Cardamine hirsuta* (Cardamineae; Gan et al. 2016), and *A. alpina* (Arabideae; Willing et al. 2015) genome annotations. The 4 diploid genomes (*A. arenosa*, *A. thaliana*, *A. alpina*, and *C. hirsuta*) show 1:3 syntenic relationships with the younger hexaploid *C. sativa* genome ( $n = 20$ ) and evolutionarily older hexaploid genome of *B. rapa* ( $n = 10$ ).

gene loss after polyploidy is common, it is not random, and the fate of duplicated genes is determined by the interplay of many forces, including dosage-induced constraints (Birchler and Veitia 2007, 2012; Conant 2014; Conant et al. 2014; Hao et al. 2022), differentially methylated transposable element (TE) density patterns among subgenomes (Hollister and Gaut 2009; Freeling et al. 2012), changes in regulatory networks (Blischak et al. 2018; Hu and Wendel 2019), and epigenetic changes after hybridization (Cheng et al. 2016; Zhao et al. 2017). These processes yield diploidized genomes with genes retained in different copy numbers and patterns of variation (Li et al. 2021). This understanding is not only significant for advancing our knowledge of plant genetics but also has implications for broader evolutionary biology.

Secondly, the knowledge gained from studying polyploidy and diploidization in Brassicales can have practical applications in agriculture. Genes retained after polyploidy harbor more genetic diversity than other genes and were important during the domestication of *Brassica rapa* (Qi et al. 2021). Therefore, a better understanding of the processes involved can lead to more efficient breeding strategies and the development of improved crop varieties.

Thirdly, resolving these struggles can open up new avenues for future research. It can enable the development of advanced tools and methods, such as POInT (the Polyploid Orthology Inference Tool), a likelihood method for modeling subgenome

partitioning and biased fractionation (Conant and Wolfe 2008), which can then be applied not only to Brassicales but also to other polyploid systems. Inferring the biased pattern of gene loss has been challenging for several reasons, including genomic rearrangement and homoeologous exchanges (Xiong et al. 2011; Mandáková et al. 2017; Edger et al. 2018b; Bird et al. 2020, 2023). Orthology inference using gene synteny can reliably differentiate subgenomes in polyploids (Lyons et al. 2008; Haug-Baltzell et al. 2017). POInT has been used to confirm that biased fractionation cannot be attributed to artifacts in synteny block inference (Conant 2020). Integrating tools like POInT and GENESPACE (Lovell et al. 2022) can incorporate synteny into phylogenetically robust models of evolution. These approaches are promising for studying the diploidization process, particularly as the wider Brassicales community generates more chromosome-scale genome assemblies from across the order.

## Epigenetics, small RNAs, and repetitive elements

### Epigenetics, small RNAs, and transposable elements across the Brassicaceae

Much of our knowledge of small RNA (sRNA) biology has been derived from mutant screens in *A. thaliana*, including



studies of TE mobilization and repression pathways (Miura et al. 2001; Gendrel et al. 2002; Lippman et al. 2004; Martinez et al. 2017). Plant genomes are typically dominated by TEs that can move throughout the genome using a variety of mechanisms (Wicker et al. 2007). Although cycles of TE turnover are crucial for shaping plant genomes, most TEs are kept in a transcriptionally repressed state via DECREASED DNA METHYLATION 1 (DDM1) and RNA-directed DNA methylation pathways (Zemach et al. 2013; Matzke and Mosher 2014; Li et al. 2015; Gallego-Bartolomé et al. 2019). The loss of DDM1 leads to loss of DNA methylation, accompanied by TE mobilization and developmental defects (Kakutani et al. 1999; Miura et al. 2001). The *A. thaliana* genome, which was the first plant genome to be sequenced (The Arabidopsis Genome Initiative 2000), has been paramount to the dissection of these pathways; however, there is immense variation in DNA methylation levels within the Brassicaceae (Seymour et al. 2014; Bewick et al. 2016), such as the loss of gene body methylation in *Eutrema salsugineum* (Eutremeae) due to a loss of the CHROMOMETHYLASE 3 gene (Bewick et al. 2016). A recent study by Lu et al. (2019) investigated epigenetic variation across 13 angiosperm species, 2 of which are in the Brassicaceae (*A. thaliana* and *E. salsugineum*). They found that species-specific accessible chromatin regions were strongly enriched for TE sequences. Therefore, species-specific TEs may regulate gene expression across the order, emphasizing the importance of sequencing across the clade.

### Epigenetics, sRNAs, and TEs across the Brassicales

In the Brassicales, sRNAs have been poorly sampled in non-Brassicaceae species, though there are several emerging studies related to classes of sRNAs. One example is research on phased, secondary, small interfering RNAs (phasiRNAs), which display an intriguing pattern of loss in the Brassicales. In *A. thaliana*, phasiRNAs are expressed in vegetative tissues (Fei et al. 2013); additionally, 21-nt and 24-nt reproductive phasiRNAs, triggered by miR2118 and miR2275, respectively, have been found broadly across monocots (Kakrana et al. 2018; Patel et al. 2018) and eudicots (Xia et al. 2019) and shown to play a role in anther fertility in corn and rice (Teng et al. 2020). Intriguingly, miR2275 is not present in any of the sampled Brassicales (Polydore et al. 2018; Xia et al. 2019). The loss of miR2275 has been reported in several other lineages, including legumes and Solanaceous species, allowing us to place the Brassicales in a larger context of sRNA loss, thus enabling comparisons across the tree of life to understand the anther fertility phenotype compared with other monocots and eudicots. Other floral-related microRNAs in the Brassicales are implicated in sex determination and sexual dimorphism in papaya (Aryal et al. 2014). Further examples of lineage-specific sRNA evolution in the Brassicales are tandem microRNA loci, miR826, and miR5090, involved in nitrogen starvation adaptation (He et al. 2014).

With multiple documented WGDs, the Brassicales are a model clade for testing the interplay between polyploidy, genome size, and repetitive element proliferation. A recent

study characterized TE abundances from low-coverage genome shotgun data in 71 phylogenetically diverse taxa across the order, placing transposon content in the context of WGD and phylogeny (Beric et al. 2021); surprisingly, little connection was found between phylogeny and TE expansion or contraction, nor was there any correlation between patterns of WGD and TE abundance. Although WGDs might be expected to lead to global TE reactivation, TE purging mechanisms seem to work efficiently in the Brassicales (Hawkins et al. 2009). Interestingly, the elimination of TEs is less effective in some Brassicales clades, resulting in large diploid genomes (1,500–4,200 Mb) that exceed the modal C-value of the Brassicales by 4- to 10-fold (Hloušková et al. 2019; Zuo et al. 2022).

### The future of Brassicales epigenetics, sRNA, and TE research

The future of epigenetics, sRNA, and repetitive element studies within the Brassicales hinges on our ability to generate diverse sRNA and methylome sequencing datasets and anchor analyses onto chromosome-scale assemblies in a comparative genomic context. Genomes from across the order will provide opportunities for investigation of how sRNA biogenesis and methylation pathways of *A. thaliana* function in phylogenetically distant relatives, as well as provide insight into polyploidy, transposon biology, and lineage-specific sRNA emergence and loss.

## Reproductive biology

### Reproductive biology across the Brassicaceae

Thirty years ago, homeotic genes discovered in *A. thaliana* formed the foundation of the “ABC model” (Bowman et al. 1991a, 1991b; Drews et al. 1991). *A. thaliana* is also a model for understanding the evolutionary genetic underpinnings of the loss of self-incompatibility (SI) which promotes outcrossing. SI is a widespread trait found in 40% of flowering plant species and is based on recognition of self-pollen (e.g. Brassicaceae, Papaveraceae), or non-self-pollen (e.g. Solanaceae; Iqic et al. 2008; Fujii et al. 2016). The mechanisms underlying sporophytic SI in the Brassicaceae were discovered in *Brassica* and involve a pollen-specific ligand (*S-LOCUS PROTEIN 11*; or *S-LOCUS CYS-RICH PROTEIN*, SCR) and pistil-specific receptor (*S LOCUS RECEPTOR KINASE*) tightly linked at the S-locus (Stein et al. 1991; Schopfer et al. 1999; Takasaki et al. 2000; Takayama et al. 2000). The loss of SI arose in *A. thaliana* at least 3 times (Shimizu et al. 2008; Tsuchimatsu et al. 2017) through loss-of-function mutations in SCR, followed by rearrangements of the S-locus (Shimizu et al. 2008; Tsuchimatsu et al. 2010, 2017). The loss of SI in the Brassicaceae has occurred independently in at least 5 other species (Shimizu and Tsuchimatsu 2015; Nasrallah 2017). Further, there is an epigenetic component to SI plasticity. In *Brassica*, both DNA methylation (Shiba et al. 2006) and 24-nt sRNAs (Tarutani et al. 2010) target recessive alleles in allelic pollen S-determinants. Both genomic and ecological factors can

influence the repeated evolution and breakdown of SI, so understanding the diversity of pathways leading to SI loss can be gained by studying systems across the Brassicales, although to date most involve the primary loss of the male-specific determinant of SI (Shimizu and Tsuchimatsu 2015).

### Reproductive biology across the Brassicales

Most species in the Brassicales are hermaphroditic or monoecious (Renner 2014). However, roughly 1% are dioecious, where unisexual flowers develop on separate plants (Renner 2014). Dioecy has evolved in 7 different families, representing at least 5 origins (Fig. 2; Renner 2014; Soza et al. 2014). Although structurally hermaphroditic, Cleomaceae commonly vary the ratio of functionally male to functionally female flowers according to environmental conditions (Zohoungbogbo et al. 2018).

In some Brassicales, sex chromosomes have been identified, with the most detailed studies in papaya (Caricaceae). The papaya Y is defined by the presence of an approximately 10-Mb nonrecombining sex-determining region that is only a fraction (approximately 10%) of the entire sex chromosome (Liu et al. 2004; Na et al. 2012; Wang et al. 2012; Yue et al. 2022); this is similar to other plant sex chromosomes like asparagus and kiwifruit (Akagi et al. 2018; Harkess et al. 2020). The closely related genus *Vasconcellea* also has evidence of XY chromosomes; however, evidence suggests this is an independent origin (Wang et al. 2012; Carvalho and Renner 2015). Although papaya is almost exclusively dioecious in the wild, it is often described as trioecious because most cultivars are essentially gynodioecious (Brown et al. 2012; Fuentes and Santamaría 2014).

The Brassicales are also an excellent model for the evolution of diverse fruit structures. The Brassicaceae fruits exhibit the greatest diversity in size and shape (Ronse De Craene and Haston 2006); however, fruit characters are highly homoplastic (Franzke et al. 2011), with clear evidence of parallel evolution of indehiscent fruits (Mühlhausen et al. 2013) and independent origins of dehiscence in Capparaceae and Cleomaceae (Hall et al. 2002). Moreover, the diverse fruit characters in the Brassicaceae play major ecological roles in fruit/seed dispersal (Sperber et al. 2017; Arshad et al. 2019; Bhattacharya et al. 2019a; Nichols et al. 2020). The recent evidence of plasticity in *Aethionema* in response to stress, with corresponding glucosinolate allocation to fruit tissues (Bhattacharya et al. 2019b), demands a holistic evo-devo analysis among other members of the Brassicales to understand the genetic and epigenetic basis of diversity in fruit characters.

### The future of Brassicales reproductive biology research

The repeated evolution of dioecy, SI, and diverse fruit types across the Brassicales makes it a powerful model order for comparative developmental and genomic analyses. For instance, are the same sterility genes or pathways involved in dioecious flower development across the Brassicales? Do

the same genes control fruit shape and dehiscence? These comparisons will be valuable because they may be potential targets for controlling sex and fruit characteristics in breeding programs for the many economically important traits found across the order. Moreover, the many origins of sex chromosomes will provide further insight into their early evolution from ancestral autosomes (Carey et al. 2021).

## Specialized metabolite diversity

### Specialized metabolite diversity across the Brassicaceae

Glucosinolates, or mustard oils, are a clade-defining characteristic of plants in the Brassicales. The presence of major ecological model systems within this group, including *Arabidopsis*, *Boechera* (Boechereae), *Brassica*, and *Streptanthus* (Thelypodieae), has allowed mechanistic ecology studies to identify the forces shaping specialized diversity in the Brassicaceae. Competition experiments between *A. thaliana* accessions from central and northern Europe in the presence of multiple herbivores showed structural variation in the glucosinolate *METHYLTHIOALKYLMALATE* (MAM) locus linked to the presence or absence of 2 different aphids across the continent (Kroymann et al. 2003; Züst et al. 2012). Recreating natural variation in isogenic lines with field trials in both *Arabidopsis* and *Boechera* showed specialized metabolism is under strong selective pressure that fluctuates locally, creating local bet-hedging (Schranz et al. 2009; Manzaneda et al. 2010; Kerwin et al. 2015, 2017). Manipulating glucosinolates within *Brassica* showed how intraspecific glucosinolate variation shaped inter- and intraspecific competition and enabled stochastic variation in species occupancy across a landscape (Lankau and Strauss 2007). Combining ecological parameters, biochemistry, and phylogenetics within *Streptanthus* has been used to understand how specialized metabolism is shaped within the family in relation to serpentine soils (Cacho et al. 2015, 2021). Interestingly, the above ecological model systems and other less studied Brassicales can co-occur within the same or similar environments. This raises the potential to develop studies investigating how specialized metabolism within one species may influence the specialized metabolism within another.

### Specialized metabolite diversity across the Brassicales

Extending the ecology work via the combination of genomics and chemistry across the entire order Brassicales has begun to illustrate how specialized metabolite pathways evolve. Glucosinolates are estimated to have first occurred at the base of the Brassicales approximately 108 MYA and span the phylogeny from the leafy greens of the Brassicaceae to the seeds of the Caricaceae, where they co-occur with the hypothesized evolutionary progenitor cyanogenic glycosides (Olafsdottir et al. 2002). Glucosinolates then diversified further approximately 43 MYA with the evolution of the aliphatic glucosinolates (Beekwilder et al. 2008; Zhang et al.



2015; Czerniawski et al. 2021). Some glucosinolates, like 2-hydroxy-2methylbutyl glucosinolate (glucocleomin), are family-specific innovations (e.g. Cleomaceae), and others like 3-benzoyloxybutyl glucosinolate (glucomalcomiin) appear to be relatively species specific (e.g. *A. thaliana*). In contrast, glucosinolates like methyl glucosinolate (glucocapparin) are found in the Cleomaceae and the Capparaceae but not in the Brassicaceae, suggesting a loss or replacement in the Brassicaceae that has yet to be characterized genetically.

Although some of these innovations in glucosinolate amino acid substrate types have been mapped on the phylogeny (Edger et al. 2018a; Kumar et al. 2019; Petersen et al. 2019), the underlying genomic mechanisms influencing these transitions have yet to be phylogenetically characterized. What is clear is that the evolution of the glucosinolate pathway has involved whole-genome, local duplication, and gene transposition events, each at different levels. In the glucosinolate pathway, whole-genome duplications gave rise to variation in the CYP79 family that determines the specific amino acid used for the glucosinolate backbone (Edger et al. 2015, 2018a) and the origin of the MAM locus derived from primary metabolism (Abrahams et al. 2020). In contrast, local structural diversity, duplications, and inversions create enzymatic variation leading to novel enzymes that create the structural modifications to the glucosinolate backbone, determining the ultimate biological activity (Kroymann et al. 2003; Hansen et al. 2008; Chan et al. 2010).

### The future of Brassicales specialized metabolite diversity research

Moving beyond glucosinolates to other specialized pathways, the identified diversity and innovations at different time scales, both within and between metabolic pathways, allow for unique comparisons using the Brassicales. For example, is there a difference in how older and younger genes evolve within a pathway and is this difference consistent across pathways? In addition to investigating how pathways are gained, the Brassicales provide the ability to study how pathways can be lost and independently recreated (Haribal et al. 2001; Rajniak et al. 2015). Additional questions related to specialized metabolism can also be addressed when sampling across the Brassicales. For example, does the evolution of cardiac glycosides within the *Erysimum* lineage (Erysimeae) influence the selective pressure on specialized metabolism in other species when they coexist or within the same species when the 2 pathways coexist (Züst et al. 2020)? Developing extensive functional, biochemical, genetic, and genomic resources within the Brassicales will allow testing these questions about how specialized metabolism is driven to novelty.

### The evolution of woodiness

#### Woodiness across the Brassicaceae

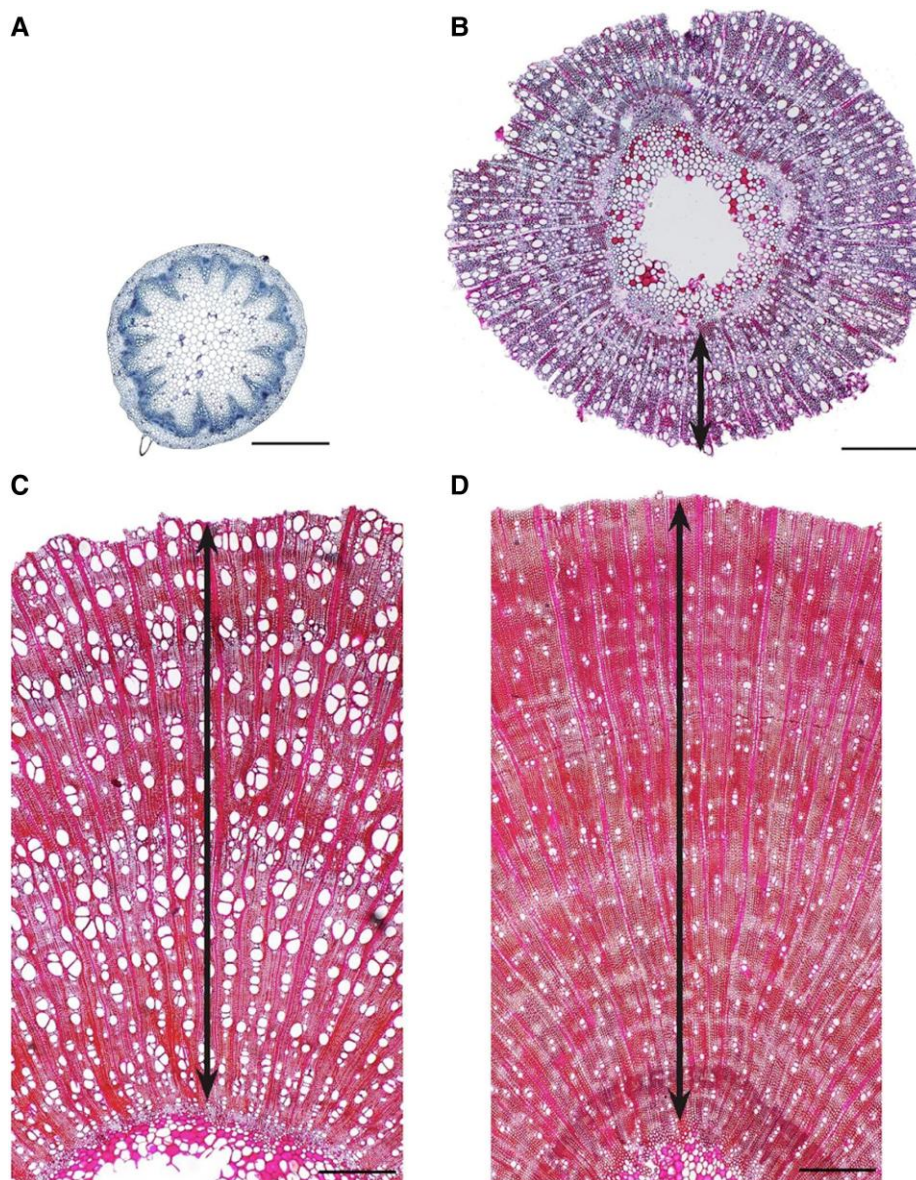
The initiation and activation of a vascular cambium can lead to massive secondary growth in woody plants, in which the

wood cylinder is the main component in stems and roots of shrubs, trees, and lianas. The herbaceous *A. thaliana* is an excellent model for elucidating the genetic controls of wood formation. Woody mutants show that the genetic mechanism turning on the wood pathway in stems can be simple, for example by knocking out 2 flowering control genes (Melzer et al. 2008; Lens et al. 2012; Davin et al. 2016) or by overexpressing a single gene that is downstream of these 2 flowering control genes (Rahimi et al. 2022). Interestingly, about 10% of Brassicaceae species have developed into woody shrubs, treelets, and even lianas under natural conditions, and these approximately 400 woody species are the result of nearly 100 independent transitions from herbaceousness toward phylogenetically derived woodiness (e.g. Mohammadin et al. 2017; Zizka et al. 2022; Lens et al., unpublished dataset). This makes the Brassicaceae a perfect model to better understand why flowering plants became woody during evolutionary history.

#### Woodiness across the Brassicales

Woodiness represents the ancestral state in the Brassicales (Figs. 2 and 4). Across the order, evolutionary transitions from ancestral woodiness toward herbaceousness and reversals back to phylogenetically derived woodiness have repeatedly occurred. Transitions toward herbaceousness evolved at least 7 times, leading to, among others, the predominantly herbaceous sister families Brassicaceae (Al-Shehbaz 1984; Franzke et al. 2011) and Cleomaceae (Byng 2014; Patchell et al. 2014) that include by far the majority of perennial or annual herbaceous species within the order. Other families including herbs are Emblingiaceae (Hall et al. 2004; Byng 2014), Gyrostemonaceae (Byng 2014), Limnanthaceae (Meyers et al. 2010; Edger et al. 2018a), Resedaceae (Martín-Bravo et al. 2007), Tovariaceae (Hall et al. 2004), and Tropaeolaceae (Andersson and Andersson 2000). Within the predominantly herbaceous Brassicales clades, we also identified approximately 100 evolutionary reversals from herbaceousness toward derived woodiness (e.g. Lens et al., unpublished dataset), accounting for about 15% of the total number of reversals within flowering plants. Nearly all of these independent reversals are in the Brassicaceae, but they also occurred in Cleomaceae (at least 4 transitions; Patchell et al. 2014) and Resedaceae (at least 3; Martín-Bravo et al. 2007).

Although the tallest Capparaceae trees reach 40 m in height, other trees in the order are typically much smaller (e.g. Akaniaceae, other Capparaceae; Bayer and Appel 2003; Mercado Gómez and Escalante 2018). Across the entire order, multiple invasions into seasonally dry (Brassicaceae, Gyrostemonaceae, Resedaceae), (hyper)arid (Brassicaceae, Cleomaceae, Gyrostemonaceae, Koerberliniaceae, Moringaceae, Salvadoraceae, Setchellanthaceae, Tiganophytaceae), and temperate habitats (Brassicaceae, Cleomaceae, Limnanthaceae) gave rise to a general reduction in plant size, leading to small shrubs and herbs in many lineages (Al-Shehbaz 1984; Martín-Bravo et al. 2007; Byng 2014;



**Figure 4.** Light microscope cross sections through Brassicales stems showing the difference between: herbaceous **A**) and woody stems **B–D**). Double-headed arrows indicate the wood cylinder. All images are at the same magnification (scale bar = 500  $\mu\text{m}$ ). **A**) Basal inflorescence stem part of *A. thaliana* (Brassicaceae). **B**) *Reseda* sp. (Resedaceae). **C**) *C. spinosa* (Capparaceae). **D**) *Sinapidendron angustifolium* (Brassicaceae).

Salariato et al. 2016; Swanepoel et al. 2020). It is noteworthy that woody Gyrostemonaceae, which may reach 10 m, remain relatively short-lived (usually <10 years) in arid areas (Baker et al. 2005). The successful attempts to establish and diversify into (hyper)arid habitats is unusual across the largely tropical rosids, making the Brassicales in general and the species-rich Brassicaceae, in particular, stand out as ecologically distinct lineages (Cornwell et al. 2014).

#### The future of Brassicales woodiness research

Ongoing efforts to start building a species-dense Brassicales phylogeny (Nikolov et al. 2019; Hendriks et al. 2023), required to identify all the species pairs with contrasting life forms, makes the Brassicales a new, representative model clade to

investigate the environmental and genetic drivers of the multiple woodiness shifts. The emerging phylogenetic framework will inform the first step toward reconstructing the evolution of niche dimensions and allow for solid statistical testing of hypotheses explaining why plants became woody (Lens et al. 2013; Dória et al. 2018; Zizka et al. 2022). Moreover, identification of the approximately 100 contrasting life form pairs in the Brassicales will serve as a great tool to help discover key regulatory genes controlling wood formation and subsequently validate these results in *A. thaliana* (Melzer et al. 2008; Lens et al. 2012; Davin et al. 2016; Rahimi et al. 2022). Using newly developed approaches such as phylogenetic inter-species genome-wide association studies (Kiefer et al. 2019), we will be able to assess whether



the same genes are involved in the independent transitions, thereby contributing to our general understanding of parallel and convergent evolution as a principal evolutionary phenomenon (Elmer and Meyer 2011).

## Photosynthesis

### Photosynthesis across the Brassicaceae

Modifications to  $C_3$  photosynthesis have repeatedly occurred in angiosperms and typically are characterized by spatial ( $C_4$ ) or temporal (crassulacean acid metabolism) sequestration of  $CO_2$  fixation (Edwards 2019). Within the Brassicaceae, so far only *Moricandia* (Brassicaceae) has been reported to include 5  $C_3$ - $C_4$  intermediate species as well as 3  $C_3$  species, whereas no species has been shown to have  $C_4$  photosynthesis (Apel et al. 1997; Schlüter et al. 2017). Schlüter et al. (2017) suggest various constraints on the path to  $C_4$ , including anatomical limitations to efficient metabolite exchange, limitation in nutrients other than carbon, and limited pressure to reduce photorespiration as these species inhabit cooler climates than those occupied by  $C_4$  species. Other Brassicaceae lineages deviate from typical  $C_3$  physiology (e.g. carbon isotope ratios not typical of  $C_3$ ), including *Diplotaxis tenuifolia* (Brassicaceae; Apel et al. 1997; Ueno et al. 2003, 2006), *Brassica gravinae* (Brassicaceae; Ueno 2011), and *Hirschfeldia incana* (Brassicaceae; Garassino et al. 2022). Such species and close relatives need detailed analysis to understand how  $C_3$  photosynthesis has been modified.

### Photosynthesis across the Brassicales

In the Brassicales, at least 4 instances of partial- or full- $C_4$  photosynthesis have evolved. In Cleomaceae,  $C_4$  NAD-dependent malic enzyme (NAD-ME) photosynthesis has been extensively characterized in *Gynandropsis gynandra* (Marshall et al. 2007; Voznesenskaya et al. 2007; Bräutigam et al. 2011; Brown et al. 2011; Koteyeva et al. 2011, 2014; Kajala et al. 2012; Williams et al. 2016; Huang et al. 2021). This research indicates that  $C_4$  photosynthesis has evolved from components found in  $C_3$  species. Identification of a bipartite transcription factor module driving gene expression in the *A. thaliana* bundle sheath provides insight into *trans*-factors important for  $C_4$  photosynthesis (Dickinson et al. 2020). Significant natural variation of  $C_4$  traits exist in *Gynandropsis* (Reeves et al. 2018), and the *G. gynandra* genome has gene families retained in duplicate following polyploidy that facilitated  $C_4$  evolution (Hoang et al. 2023). The *Coalisina* clade (Cleomaceae; Roalson and Hall 2017) includes  $C_3$ , intermediate, and  $C_4$  species (Voznesenskaya et al. 2007). *Coalisina angustifolia* has NAD-ME  $C_4$ , whereas *Coalisina paradoxo* has been characterized as intermediate/ $C_2$  (Koteyeva et al. 2011). Only 1 taxon within the *C. angustifolia* complex has been fully characterized (*C. angustifolia* subsp. *angustifolia*; Voznesenskaya et al. 2007). Denser sampling of this clade would inform us about the  $C_3$  to  $C_4$  transition as it appears to include  $C_3$ ,  $C_3$ - $C_4$  intermediate, and fully  $C_4$  species

and would provide an interesting comparative lineage to *Moricandia*, where there are  $C_3$  and  $C_3$ - $C_4$  intermediates but no known  $C_4$  species. *Areocleome* is the least studied of Cleomaceae  $C_4$  lineages and includes at least 1 species restricted to arid Australia (Barrett et al. 2017). *Areocleome oxalidea* has NAD-ME-type  $C_4$  physiology (Voznesenskaya et al. 2007; Koteyeva et al. 2011), but the underlying  $C_4$  genetics remain unstudied. The sister genus *Arivela* may have  $C_3$ - $C_4$  intermediate characteristics as it is sympatric with *Areocleome* and more widespread in the monsoonal tropics. Other Cleomaceae lineages, including *Cleomella sparsifolia*, *Tarenaya siliculifera*, *Sieruela allamanii*, and *S. gallaensis*, appear to deviate from typical  $C_3$  physiology (Voznesenskaya et al. 2007; Parma et al. 2021). These species significantly differ in their distributions from dry southwestern North America to tropical Brazil, and tropical to subtropical Africa, providing opportunities to consider  $C_3$  pathway modifications across environments.

### The future of Brassicales photosynthesis research

Understanding the modifications in photosynthetic pathways and carbon-concentrating mechanisms holds significant relevance and impact in several key aspects. Gaining insights into these fundamental physiological processes is crucial for advancing our comprehension of plant biology and evolution. This knowledge helps us unravel the intricacies of how plants have adapted to various environmental conditions, contributing to a deeper understanding of the natural world.

Photosynthetic pathway modifications occur in numerous lineages and in some cases co-occur and shift between the 2 major modification types (Holtum et al. 2017; Edwards 2019; Winter et al. 2019). Further,  $C_4$  photosynthesis has multiple biochemical forms (Sage 2004). As our understanding of these basic physiological processes has expanded, 4 foundational hypotheses have solidified: (1) there have been multiple independent origins of  $C_4$ /CAM (Sage et al. 2011); (2) independent origins of carbon-concentrating mechanisms can occur through the co-option of similar molecular components (Christin et al. 2007; Brown et al. 2011), and these components exist in the ancestral  $C_3$  state (Brown et al. 2011; Williams et al. 2016; Reyna-Llorens et al. 2018); (3) precursor anatomical modifications that appear unrelated to photosynthesis act as facilitators of biochemical adaptations (Christin et al. 2013; Williams et al. 2013); and (4) a number of carbon-concentrating mechanisms do not fully fit our concepts of  $C_4$ /CAM (e.g.  $C_2$ ; Lundgren 2020). Our understanding of carbon-concentrating adaptations has become better attuned to these nuances as knowledge on independent origins has accumulated.

Resolving the current struggles in this field would open up exciting future directions for research. Access to genomes and a comprehensive understanding of photosynthetic pathway modifications would enable researchers to explore the genetic and molecular underpinnings of these processes in even greater detail. This, in turn, could lead to the discovery



of novel molecular components and mechanisms that drive photosynthetic adaptations. The relevance of this research extends to agriculture and environmental conservation. By understanding the multiple independent origins of  $C_4$ /CAM photosynthesis and the co-option of molecular components, we can potentially engineer crops to be more efficient in photosynthesis and better adapted to changing environmental conditions. This has implications for food security and sustainable agriculture, particularly in the face of climate change. Additionally, the recognition of carbon concentrating mechanisms that do not fit traditional  $C_4$ /CAM concepts, such as  $C_2$ , presents an intriguing avenue for further investigation. Unraveling the molecular basis of these mechanisms could challenge and expand our current understanding of photosynthesis and carbon fixation in plants.

## Extremophytism and stress tolerance

### Extremophytism across the Brassicaceae

Many Brassicaceae (and Brassicales) species thrive in extreme environments including desert, arctic, and saline ecosystems. To survive in such habitats, extremophytes possess genetic adaptations for tolerating severe abiotic stresses. Such adaptations are exemplified in 2 halophytic Brassicaceae, *Schrenkiella parvula* (Schrenkielleae) and *Eutrema salsugineum* (Eutremeae), quintessential models (Zhu 2015) for studying tolerance to ionic and nutrient stresses (Kazachkova et al. 2018; Pantha et al. 2021; Tran et al. 2021; Wang et al. 2021). Curated genomes, multiple tissues, and environmental response “omics” datasets have been generated for both species as well as transformation protocols that facilitate independent genetic analyses (Oh et al. 2010, 2014; Dassanayake et al. 2011; Wu et al. 2012; Wang et al. 2019; Pantha et al. 2021; Tran et al. 2022; Wijesinghege et al. 2022a, 2022b; Sun et al. 2022).

Because multiple evolutionary paths can provide unique solutions to different environmental challenges, the Brassicaceae from various extreme environments are being studied including: (1) the desert species *Anastatica hierochuntica* (Anastaticaceae), which is tolerant to heat, low nutrient, and salt stress (Eshel et al. 2017, 2022); (2) the metal hyperaccumulator *Arabidopsis halleri* (Arabidopsidae; Briskine et al. 2017; Honjo and Kudoh 2019); and (3) 3 Arctic species, *Cardamine bellidifolia* (Cardamineae), *Cochlearia groenlandica* (Cochlearieae), and *Draba nivalis* (Arabideae; Birkeland et al. 2020). Together, these models should provide a window into evolutionary adaptations that confer stress tolerance and facilitate an extremophyte lifestyle.

### Extremophytism across the Brassicales

Beyond the Brassicaceae, only a few studies have examined extremophyte Brassicales. For instance, *Batis maritima* (Bataceae), a perennial, succulent halophytic shrub that inhabits mangrove swamps, salt marshes, and salt flats (Marcone 2003; Debez et al. 2010), tolerates up to  $1\text{ M NaCl}$  and has

evolved different salt tolerance mechanisms compared with the 2 annual Brassicaceae halophyte models, *E. salsugineum* and *S. parvula*. For example, *B. maritima* accumulates high levels of shoot  $\text{Na}^+$  while *E. salsugineum* minimizes entry of  $\text{Na}^+$  (Kant et al. 2006; Debez et al. 2010).

Drought responses were investigated in the stress-tolerant, evergreen, African tree species, *Dobera glabra* (Salvadoraceae). When compared with co-occurring non-Brassicaceae species, *D. glabra* displayed the lowest water potential, illustrating its ability to thrive in drought-prone areas (Gebrekirstos et al. 2014).

Molecular studies of non-Brassicaceae Brassicales are extremely sparse. Mercati et al. (2019) reported a de novo transcriptome assembly from the drought-tolerant, xerophytic crop, *Capparis spinosa* (Capparaceae), and they generated the first set of SSR markers distinguishing subspecies of *C. spinosa*. A chromosome-level *C. spinosa* var. *herbacea* genome assembly showed expansion of gene families involved in photosynthesis and response to abscisic acid (Wang et al. 2021). Phylogenetic analysis of a Brassicales-specific gene encoding *PROTEIN PHOSPHATASE7-LIKE (PP7L)* involved in chloroplast development and abiotic stress tolerance suggested that *PP7L* evolved about 43–92 MYA, after the divergence of Caricaceae from the other Brassicales families but before the separation of Cleomaceae from Brassicaceae (Xu et al. 2019).

### The future of Brassicales extremophyte research

Core stress response pathways exist in all land plants. Yet, selective pressures imposed by extreme environments have driven novel evolutionary innovations to modify these core pathways or add alternatives. These evolutionary innovations are being revealed by comparative analyses of the premier model species *A. thaliana* (“stress-sensitive”) with the Brassicaceae extremophyte models. Yet these Brassicaceae species are all annuals, and alternative stress tolerance strategies could be found in perennial extremophytes that survive harsh stresses over multiple years. Thus, the gap in the adaptive trait space left by the Brassicaceae models can be narrowed by studies of perennial Brassicales extremophytes, which possess the added advantage of shared ancestry with *A. thaliana*, many of whose core stress response pathways have been identified.

The lack of genomic resources for the Brassicales extremophytes has limited research into their adaptations to extreme environments. Emerging technologies in large-scale and single-molecule/cell sequencing offer unprecedented molecular tools that require little prior genetic information to produce primary genomic resources. Single-molecule/hyperspectral imaging and phenomics can then facilitate linkage of novel genomic variation to unique extremophyte traits. Thus, the broader phylogenetic footprint capturing the life history strategies of the Brassicales extremophytes could facilitate new discoveries for sustainable crop production, land reclamation, and biodiversity conservation in this

era of climate change and global threats to the environment and food insecurity.

## Supporting model orders with global communities

Issues identified not only in the determination of relationships among or within the Brassicales but also in analysis of all the phenotypes discussed here create many opportunities to leverage the trait diversity across the order. To fully take advantage of these opportunities for both those currently investigating this diverse plant order and those who wish to, there must be a cohesive development of a global network for plant material, data generated, and researchers. Turner-Hissong et al. (2020) highlight ways to curate community efforts in organizing not only germplasm and data but also outreach and broader impacts. Although these suggestions are primarily directed at crop systems, many ideas are useful here too. For example, there is a need to develop reliable stock sources or live genetic material, which researchers can both order and submit samples, to increase reproducibility across studies. Some examples of this exist, such as the *Arabidopsis* Biological Resource Center (ABRC; <https://abr.csu.edu/>), the Universidad Politécnica de Madrid Plant Germplasm Bank (UPM-PGB; which includes the César Gómez Campo Collection), the USDA Germplasm Resources Information Network (GRIN), and the Royal Botanic Gardens (Kew) Millennium Seed Bank and its DNA and tissue bank collections. However, one must first know about these resources to utilize them. However, even with these resources available, cultivating them demands substantial time, effort, funding, and space. Numerous species lack established protocols, particularly those accessible to the public, on optimizing plant growth for generating sufficient seeds for trait-focused studies. It is essential to meticulously document the origins of genetic resources and acknowledge the knowledge's source by returning it to the country of origin.

Following germplasm curation, data curation of all types is crucial for continual progress. As we generate these large, multi-omic datasets, there is an increasing need for scalable, systematic community efforts to organize these resources, all of which will require standards for data collection and curation and ideally utilize established open-access repositories. Fortunately, there are already databases in existence, such as BrassiBase (Koch et al. 2018), which consolidate information encompassing taxonomy, systematics, evolution, available germplasm resources, and specimen collections and could be expanded to include all of the Brassicales in the future. Furthermore, networks like the Multinational *Brassica* Genome Project (MBGP; [brassica.info](http://brassica.info)) could play a pivotal role if expanded to Brassicales. Currently, MBGP organizes annual meetings at international conferences and is dedicated to establishing a *Brassica* Information System, which includes the formulation of standards for describing data

entities and experimental resources. Additionally, platforms like Phytozome, which serves as a central hub for accessing, visualizing, and analyzing JGI-sequenced plant genomes, along with the recently published and updated JGI Gene Atlas (Sreedasyam et al. 2023), can greatly contribute to the development of a cohesive community focused on the Brassicales model clade. These types of databases provide a place where not only data can be accessed, but researchers can also keep others updated on current projects, results, and future plans. Centralizing access to these plant materials and data generated would significantly decrease the barriers for scientists, especially early career scientists and those with less access to modern resources, and ultimately encourage equity and recruit new and diverse researchers to study the model order Brassicales. Nonetheless, persistent challenges exist, including a decline in taxonomic expertise, restricted access to materials from several small endemic families, and a noticeable bias toward collaboration with researchers primarily from the global North.

Greater accessibility to plant genomics at lower costs has expanded global collaboration opportunities. However, it is important to acknowledge that the discovery of genes and pathways in plant species can have significant economic implications, so these initiatives must also address the growing complexity surrounding genetic resources and intellectual property rights. There are growing calls for decolonization of science in the global North that extracts materials and knowledge from the global South and Indigenous groups without reciprocity (Radcliffe 2017; Armstrong and McAlvey 2019; Baker et al. 2019; Carroll et al. 2020; The First Nations Information Governance Centre 2022). Within the order Brassicales, *Pentadiplandra brazzeana* was at the center of a biopiracy controversy when scientists from the University of Wisconsin patented a protein, brazzein, extracted from the berries of Gabonese plants as a sweetening agent with no benefit-sharing agreement with Gabon or Gabonese people (Blakeney 2019; Dwivedy et al. 2019). A growing number of countries have implemented the Nagoya Protocol on Access and Benefit-sharing (Buck and Hamilton 2011) and other measures to govern the exchange and use of biological materials. Moving forward, we urge all researchers working on Brassicales collection and sequencing efforts to consider genuine collaboration with Indigenous groups and scientists in the global South to work to align project goals with local interests (Baker et al. 2019). We understand the need to build a foundation of respect, reciprocity, and accountability within our growing collaborative efforts in keeping with the principles of Indigenous data sovereignty and equitable benefit sharing. The authors here commit to these core principles throughout our contacts and research.

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## Author contributions

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