Origin and Evolution of Diploid and Allopolyploid Camelina Genomes Were Accompanied by Chromosome Shattering

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Complexes of diploid and polyploid species have formed frequently during the evolution of land plants. In false flax (Camelina sativa), an important hexaploid oilseed crop closely related to Arabidopsis (Arabidopsis thaliana), the putative parental species as well as the origin of other Camelina species remained unknown. By using bacterial artificial chromosome–based chromosome painting, genomic in situ hybridization, and multi-gene phylogenetics, we aimed to elucidate the origin and evolution of the polyploid complex. Genomes of diploid camelinas (Camelina hispida, n = 7; Camelina laxa, n = 6; and Camelina neglecta, n = 6) originated from an ancestral n = 7 genome. The allotetraploid genome of Camelina rumelica (n = 13, N6H) arose from hybridization between diploids related to C. neglecta (n = 6, N6) and C. hispida (n = 7, H), and the N subgenome has undergone a substantial post-polyploid fractionation. The allohexaploid genomes of C. sativa and Camelina microcarpa (n = 20, N6N7H) originated through hybridization between an auto-allotetraploid C. neglecta–like genome (n = 13, N6N7) and C. hispida (n = 7, H), and the three subgenomes have remained stable overall since the genome merger. Remarkably, the ancestral and diploid Camelina genomes were shaped by complex chromosomal rearrangements, resembling those associated with human disorders and resulting in the origin of genome-specific shattered chromosomes.

INTRODUCTION

Polyploidization is an important speciation mechanism, particularly in ferns and angiosperms. New polyploid species form repeatedly, and go extinct or return to the genetically diploid state via a process described as (re)polyploidization. Assuming that early angiosperm divergence was predated by a whole-genome duplication (Jiao et al., 2011), the polyploidization–diploidization cycles mark speciation and cladesogenesis of angiosperms for the past ~300 million years (Clark and Donoghue, 2017).

When encountering polyploidy in extant plant genomes, two key factors immediately emerge: the way in which polyploids are formed and their age. Both autopolyploidy (genome doubling within a species) and allopolyploidy (genome merger due to interspecies hybridization) gave rise to new clades and species, many of which thrived and still exist today (Parisod et al., 2010; Jiao et al., 2011; Garsmeur et al., 2014; Marcussen et al., 2014; Landis et al., 2018). The merging of reduced and unreduced gametes in polyploids forms triploids or allotetraploids, and additional polyploid genomes can arise by backcrossing to the diploids or hybridization with other congeneric polyploids. This network of parental and hybrid-derived species of different ploidy represents a polyploid complex (Stebbins, 1971).

Polyploid complexes exhibit different levels of completeness and maturity (Stebbins, 1971) as genomes of diploid progenitors, as well as polyploid species, further evolve or eventually become extinct. The continuous extinction and genome evolution including rediploidization hamper the identification of parental (sub) genomes and inter-genome comparisons. A number of model and crop polyploid complexes, such as Brassica (Nagaharu and Nagaharu, 1935; Wang et al., 2011; Yang et al., 2016), Capsella (Douglas et al., 2015), Cardamine (Mandáková et al., 2013, 2019), Fragaria (Edger et al., 2019), Gossypium (Paterson et al., 2012; Hu et al., 2019), Nicotiana (Renny-Byfield et al., 2013; Sierro et al., 2014), Spartina (Salmon et al., 2005), Tragopogon (Symonds et al., 2010), and Triticum (Marcussen et al., 2014; El Baidouri et al., 2017), have been investigated over the years. Significant progress in high-throughput whole-genome sequencing of these and other diploid and polyploid genomes has greatly advanced our understanding of post-polyploid genome evolution. Despite recent technological advances, analysis of complex polyploid genomes remains challenging (Kyriakidou et al., 2018), especially when the parental genomes are extinct or cannot be easily identified.

Among crucifer species (Brassicaceae), Brassica is the most researched polyploid complex composed of three main diploid (2n = 16, 18, and 20; BB, CC, and AA genome) and three allo-tetraploid (2n = 34, 36, and 38; BBCC, AABB, and AACC) genomes (Nagaharu and Nagaharu, 1935). Because of its worldwide economic importance, the genus Brassica has become one of the more important models in plant polyploidy research (Mason and Snowden, 2016; Gaebelien and Mason, 2018). Recently, false flax (Camelina sativa), an increasingly popular crucifer oilseed crop, was suggested to have an allohexaploid origin (Hutcheon et al., 2010; Kagale et al., 2014), similar to rediploidized
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Background: Camelina (Camelina sativa; gold-of-pleasure or false flax) is an ancient oilseed crop grown in Europe as early as 4000 BC. Gold of pleasure was largely forgotten as a crop, being replaced by oilseed rape, and recently re-discovered for its unique characteristics. The high oil content (28–40%) makes camelina an important aviation biofuel, while its omega-3 fatty acids (“fish oils”) are beneficial for cardiovascular health. Sequencing of the camelina genome revealed that the genome consists of three parental subgenomes. However, how the hexaploid camelina genome originated and how the three subgenomes are related to other genomes in the genus Camelina remained unknown. We tackled these questions by using comparative chromosome painting, genomic in situ hybridization, and multi-gene phylogenetic analyses.

Question: We knew that the C. sativa genome consists of three subgenomes merged through a hybridization between older parental species. We wanted to find the most probable parental genomes of the hexaploid camelina and moreover to understand how its genome is related to genomes of other Camelina species.

Findings: Genomes of diploid camelinas (C. hispida, n = 7 chromosomes; C. laxa, n = 6; and C. neglecta, n = 6) originated from an ancestral n = 7 genome. The allotetraploid C. rumelica genome (n = 13, NFL genome) arose from hybridization between diploids related to C. neglecta (n = 6, NFL) and C. hispida (n = 7, H), and the N subgenome has been substantially reshuffled by chromosomal rearrangements. The allohexaploid genomes of C. sativa (n = 20, NFLH genome) originated through hybridization between an auto-allotetraploid C. neglecta-like genome (n = 13, NFLH) and C. hispida (n = 7, H), and the three subgenomes remained overall stable since the genome merger. Remarkably, the ancestral and diploid Camelina genomes were shaped by complex chromosome shattering, resembling similar events associated with human disorders.

Next steps: There are multiple implications of our work. Breeders want to improve the seed yield, oil content and other agronomic traits of the camelina crop. This can be done by producing artificial camelina lines by crossing the now identified parental Camelina species. The diploid and tetraploid Camelina genomes should be more amenable to genome sequencing and uncovering the key metabolic pathways. Future work should focus on elucidating genomic homogeneity of Camelina species analyzed in our study.

Brassica genomes. Although both Brassica and Camelina polyploid complexes include allohexaploid genomes and genomic resources available for both genera allow for insightful comparisons, the extant as well as ancestral Camelina genomes remain virtually unknown.

The genus Camelina, represented by seven or eight species (Brock et al., 2019), belongs to one of the most karyologically variable crucifer genera, with chromosome numbers ranging from 2n = 12 to 40 (2n = 12, 14, 16, 26, 28, 32, 36, 38, and 40) and a threefold genome size variation (BrassiBase, https://brassibase.cos.uni-heidelberg.de/; Hutcheon et al., 2010; Brock et al., 2018). C. sativa (2n = 40), is an ancient oilseed crop with a newfound application as an aviation biofuel and omega-3-rich feedstock. Two species, Camelina microcarpa (2n = 40) and Camelina rumelica (2n = 26), are worldwide weeds. Three other species, Camelina hispida (2n = 14), Camelina laxa (2n = 12), and Camelina neglecta (2n = 12), occur in relatively restricted ranges in Europe and western Asia. Because of variable chromosome numbers in diploids (2n = 12 and 14) and uncorrelated numbers in polyploids (2n = 26 and 40), the evolution of the Camelina polyploid complex was a conundrum until recently. Hutcheon et al. (2010) revealed that three single-copy nuclear genes are present as three paralogous copies in the genome of C. sativa and suggested that the species most likely had an allohexaploid origin. Later genome sequencing of the C. sativa genome corroborated its allohexaploid origin by revealing three minimally diverged subgenomes of an unknown identity and origin (Kagale et al., 2014).

Recent developments on plant-based biofuels have spurred an increased interest in C. sativa as an oilseed crop. With high levels of long-chain hydrocarbons, C. sativa oil is well suited as an aviation biofuel, which has proven efficacious in jet aircraft with a 75% reduction in CO2 emissions over traditional petroleum jet fuels (Shonnard et al., 2010). Seed oil content in C. sativa ranges between 36 and 47% and has oil yields of 540 to 1410 kg/ha, which is higher than soybean (Glycine max) oil yield, but comparable to that of rapeseed (Brassica napus; Moser, 2012). Moreover, genetic resources for C. sativa have grown considerably in recent years and include efficient transformation protocols (Lu and Kang, 2008; Ruiz-Lopez et al., 2014), an available reference genome (Kagale et al., 2014), molecular genetic characterizations (Vollmann et al., 2005; Gehringer et al., 2006), and CRISPR/Cas-based technology. Through genome editing, C. sativa lines have been produced with increased oleic acid (Jiang et al., 2017; Morineau et al., 2017), decreased very long chain fatty acids (Ozseyhan et al., 2018), and increased oil content (YIELD10; Waltz, 2018). Furthermore, its close phylogenetic relationship to Arabidopsis (both Arabidopsis and Camelina are classified as members of the tribe Camelineae; Beilstein et al., 2006, 2008; Nikolov et al., 2019) gives C. sativa a unique ability to benefit from the ongoing developments in the world’s most well-studied plant model.

Given C. sativa’s rising importance as a plant model for biofuel development (Iskandarov et al., 2014), high-value molecule factory (Augustin et al., 2015; Iven et al., 2016; Augustin et al., 2017), and omega-3-rich feedstock, surprisingly little is known about its evolutionary history and subgenome origins. Little genetic diversity existing in currently available C. sativa cultivars (Vollmann et al., 2005; Brock et al., 2018; Luo et al., 2019) limits the effectiveness of traditional breeding programs. To this end, the identification of diploid progenitors, wild relatives, and the elucidation of hybridization histories are necessary requisites for further improvement of Camelina genomes for seed oil production.
Herein, we aimed to reconstruct the origin of the allohexaploid genome of *C. sativa* and elucidate the identity of its three subgenomes (Kagale et al., 2014) by comparative chromosome painting (CCP), genomic in situ hybridization (GISH), and a set of nuclear gene markers. By including all known diploid and polyploid *Camelina* species, we expected to (1) identify probable parental genomes of *C. sativa* and other *Camelina* polyploids, (2) elucidate evolutionary relationships among all *Camelina* genomes and species, (3) analyze subgenome (in)stability in the polyploid genomes, and (4) uncover mechanisms driving genome evolution in *Camelina*.

**RESULTS**

*Camelina* Diploids Exhibit Highly Reshuffled Genome Structures

All analyzed plants of *C. hispida* had seven chromosome pairs (*n* = 7), whereas six pairs were counted in *C. laxa* and *C. neglecta* (*n* = 6; Figure 1). Since the ancestral crucifer karyotype (ACK, *n* = 8, AK1 to AK8; Figure 1; Supplemental Figure 1, Lysak et al., 2016) has been considered the ancestral genome of *Camelina* and the tribe Camelinae, genome structures of the investigated *Camelina* species are described here in relation to the eight chromosomes and 22 genomic blocks (GBs; Supplemental Data Set 1A) of ACK. In the analyzed diploid species, the used chromosome-specific Arabidopsis bacterial artificial chromosome (BAC) contigs, unambiguously hybridized either to one pachytene bivalent or one pair of mitotic metaphase chromosomes. In *C. hispida*, three of seven chromosome pairs (Ch1, Ch3, and Ch7) retain the ancestral structure as in ACK (AK1, AK3, and AK7, respectively). Chromosomes Ch2 and Ch4 show highly reshuffled combinations of GBs contributed by three different ancestral chromosomes (AK2, AK4, and AK5). Chromosomes Ch5 and Ch6 resemble ancestral chromosomes AK6 and AK8 modified by a whole-arm translocation (Figure 1; Supplemental Data Set 1B). Three chromosomes in *C. neglecta* (Cn1, Cn3, and Cn5) have the ancestral-like structure of AK1, AK3, and AK6, respectively. The remaining three chromosomes combine blocks of two (Cn4 and Cn6) or three ancestral chromosomes (Cn2), respectively (Figure 1; Supplemental Data Set 1C). In *C. laxa*, only chromosome Cl1 resembles the ancestral chromosome (AK1), whereas the remaining chromosomes consist of reshuffled blocks from two (Cl4 and Cl6), three (Cl3 and Cl5), or four (Cl2) ancestral chromosomes (Figure 1; Supplemental Data Set 1D).

The Origin of the Ancestral *Camelina* Genome

The ancestral *Camelina* genome (CAM; *n* = 7, chromosomes CAM1 to CAM7; Figures 1 and 2; Supplemental Data Set 1E) has been inferred based on the following premises: (1) it has descended from the older ACK; (2) its chromosome number equalled *n* = 7, as most present-day *Camelina* genomes are based on or were derived from seven chromosomes; and (3) two CAM chromosomes (CAM2 and CAM4), combining GBs of ancestral chromosomes AK2, AK4, and AK5, were identified in all *Camelina* diploids.

Chromosomes CAM1, CAM3, CAM5, CAM6, and CAM7 structurally mirror chromosomes AK1, AK3, AK6, AK8, and AK7 of ACK, respectively. Chromosome CAM4 likely originated by a reciprocal translocation involving a major part of the bottom arm of AK4 (~5.1 Mb in the *A. thaliana* genome; breakpoint within block J, in BAC clone T14G11 [AC002341]) and a minor part of the bottom arm of chromosome AK5 (~1.53 Mb in GB M-N; breakpoint between BACs F17J16 (AL163527)/F25L23 (AL356014)). The resulting translocation chromosome consists of AK5-derived GBs K-L and M-Na, and block Jb contributed by AK4. The second translocation product, bearing blocks I, Ja, and M-Nb, fused with chromosome AK2. The resulting CAM2 chromosome underwent a complex chromosome shattering (i.e., several subsequent translocations, peri- and paracentric inversions) and bears highly reshuffled blocks D (Da, Db) and E (Ea, Eb, Ec) from AK2, I and Ja from AK4, and M-Nb from AK5.

Evolution of CAM during the Divergence of Diploid *Camelina* Species

The parsimoniously inferred structure of the ancestral *Camelina* genome allowed us to reconstruct how the modern diploid genomes originated (Figures 1 and 2; Supplemental Figures 2 to 5; Supplemental Data Sets 1B to 1D). In *C. hispida*, chromosomes Ch5 and Ch6 originated by a whole-arm reciprocal translocation involving the upper arms of CAM5 (~3.94 Mb, blocks O and P) and CAM6 (~2.55 Mb, block V), with both breakpoints in the (peri) centromeric regions. The resulting translocation chromosomes Ch5 and Ch6 consist of GBs V, Q, and R and O and P, W, and X, respectively. Chromosomes Ch2 and Ch4 consist of blocks J, Db, Ea, and Nb and K-L, M-Na, Eb, I, Ec, and Da, respectively (Figure 3). These chromosomes originated by a reciprocal translocation between CAM2 (~7.76 Mb, blocks Eb, I, Ec, and Da) and CAM4 (~5.1 Mb, block Jb), involving breakpoints Ja/Eb [T14G11 (AC002341)/T6C23 (AC013289)] and M-Na/Jb [F17J16 (AL163527)/T14G11 (AC002341)]. The breakpoints Ja/Eb and M-Na/Jb can be considered as evolutionarily re-used as they previously played a role in the origin of chromosomes CAM2 and CAM4 from AK2, AK4, and AK5 (Supplemental Data Set 1E).

In *C. neglecta*, chromosome number reduction from *n* = 7 to *n* = 6 was mediated by an end-to-end translocation between chromosomes CAM6 and CAM7 (Figures 1 and 2; Supplemental Figure 3; Supplemental Data Set 1C). Breakpoints occurred in the (sub)telomeric regions of the upper arms of both chromosomes, and the end-to-end translocation event led to the origin of the fusion chromosome Cn6 (~21.37 Mb; blocks U, T, S, V, W, and X). The Cn6 centromere came from CAM5, whereas the CAM6 centromere was eliminated during/after the fusion. Chromosome Cn2 is homoeologous to CAM2; however, its structure was altered by massive genome shattering (Figure 3).

Unlike *C. neglecta*, descending dysploidy from *n* = 7 to *n* = 6 in *C. laxa* was not mediated by a single end-to-end translocation. The genome of *C. laxa* (Figures 1 and 2) underwent a complex five-step rearrangement involving four ancestral chromosomes (CAM3, CAM5, CAM6, and CAM7; Supplemental Figures 4 and 5; Supplemental Data Set 1D): (1) An end-to-end translocation between CAM3 and CAM5, with breakpoints in the (sub)telomeres of the upper and bottom arm, respectively, was accompanied by the
Figure 1. Comparative Genome Structures of Camelina Diploids.

(A) Circos graphic displaying chromosomal collinearity between ACK (n = 8; innermost circle), CAM (n = 7), C. hispida (n = 7), C. neglecta (n = 6), and C. laxa (n = 6). Color coding and capital letters (A to X) correspond to eight chromosomes and 22 GBs of the ACK, respectively. See Figure 2 and Supplemental Data Sets 1A to 1E for boundaries of GBs.
elimination of the CAM5 centromere. (2) A whole-arm translocation between the upper arms of the fusion chromosome CAM3/5 (−10.39 Mb, blocks F and G) and chromosome CAM6 (−2.5 Mb, block VI). (3) A reciprocal translocation between CAM3/6 [−6.45 Mb, block Fa, breakpoint between BACs K24M9 (AP001303) and MVE11 (AB026654)] and CAM7 [−3.14 Mb, block Ub, breakpoint between BACs F11C18 (AL049607) and F10N7 (AL021636)]. (4) Chromosome CAM3/5/6 was altered by an −3.42-Mb pericentric inversion involving block H and became C. laxa chromosome Cl5. (5) Chromosome CAM3/6/7 lost an −610-kb segment (Wa) through an unequal translocation with Cl2 and became chromosome Cl3. Chromosome Cl2 is homeologous to CAM2, however its structure was altered by massive chromosome shattering (Figure 3).

**Genome Structure and Origin of the Allotetraploid C. rumelica**

All the analyzed plants of *C. rumelica* had 26 chromosomes (*n* = 13; Supplemental Figure 6). The unambiguous identification of all GBs in four copies on mitotic chromosomes and in two copies on pachytene spreads, confirmed the assumed tetraploid origin of the species. To elucidate the parentage of *C. rumelica*, we performed GISH with differently combined genomic DNAs (gDNAs) of the diploid species. Only gDNA of *C. h ispida* and *C. neglecta* hybridized to 14 and 12 chromosomes, respectively (Supplemental Figure 6). Thus, GISH strongly suggested an allotetraploid origin of *C. rumelica* (*2n = 26, N6N6HH*) from hybridization between genomes related to *C. hispida* (*2n = 14, HH*) and *C. neglecta* (*2n = 12, N6N6*).

Combining GISH and CCP, chromosomes of the two subgenomes were unequivocally identified: chromosomes Cr1 to Cr6 were assigned to the N6 subgenome (*C. neglecta*, *n* = 6), whereas chromosomes Cr7 to Cr13 belong to the H subgenome (*C. hispida*, *n* = 7). The absence of inter-subgenomic chromosomal translocations points to subgenome stability since the allopolyploidization event.

All but one (Cr1) chromosome in the N6 subgenome are highly reshuffled. The origin of chromosome Cr2 can be reconstructed by inferring a paracentric inversion (−10.98 Mb, blocks GBs Ja, Eb, I, Ec, and Da) followed by a pericentric inversion (−6.67 Mb, blocks Ea and D) on the ancestral CAM2 chromosomes (Supplemental Figure 7). This suggests that the origin of the allotetraploid genome of *C. rumelica* occurred prior to chromosome shattering forming the modern chromosome Cn2 in *C. neglecta* (Supplemental Figure 3). Although the N6 subgenome chromosomes Cr3, Cr4, Cr5, and Cr6 are reshuffled with complex *C. rumelica*-specific translocations and inversions, associations of GBs specific for the *C. neglecta* fusion chromosome Cn6 (CAM6 + CAM7) are still detectable within chromosomes Cr4 (blocks Ua+Xc) and Cr5 (blocks S+V+Wa+Ub+Xb; Supplemental Figures 6 and 7; Supplemental Data Set 1F).

Except for chromosomes Cr8 and Cr10, the remaining five chromosomes of the H subgenome are structurally identical with chromosomes of *C. hispida*, including the *C. hispida*-specific translocation chromosomes Ch5 (Cr11) and Ch6 (Cr12; Supplemental Figures 6 and 7; Supplemental Data Set 1F). Chromosomes Cr8 and Cr10 do not exhibit *C. hispida*-specific association of GBs (Supplemental Figure 2). Instead, the chromosomes have the ancestral structure of CAM2 and CAM4, modified by a single paracentric inversion on Cr8 (−10.98 Mb, blocks Ja, Eb, I, Ec, and Da), which was identical in both subgenomes (compare with Cr2 in the N6 subgenome). These results suggest that chromosomes Ch2 and Ch4 in the parental *C. hispida*-like genome were shuffled only after the origin of *C. rumelica*.

**C. sativa and C. microcarpa: Allohexaploid Origin and Prevailing Subgenome Stasis**

All the analyzed plants of *C. sativa* and *C. microcarpa* had 40 chromosomes (*n* = 20; Figure 4). The identification of all GBs in 6 and 12 copies on meiotic (pachytene) and mitotic chromosomes, respectively, confirmed the hexaploid status of both genomes. The gDNA probes of *C. hispida* (HH) and *C. neglecta* (NN) hybridized to 14 (HH) and 26 (NNNN) chromosomes, respectively, in both *C. sativa* and *C. microcarpa* (Figure 4). No inter-subgenomic translocations were identified by GISH, suggesting that the parental subgenomes remained stable since the allopolyploidization event. GISH results provide convincing evidence that the allohexaploid *C. sativa* and *C. microcarpa* (N6N6HN6H) genomes originated from hybridization between a not yet identified tetraploid *C. neglecta*-like genome (N6N6N6N6; subgenomes G1 and G2) and the diploid *C. hispida*-like genome (HH; subgenome G3). The alternative assignment of genomes N6 and N7 as subgenomes G1 and G2, and the H genome as subgenome G3, follows the designations introduced by Kagale et al. (2014).

The structure of the *C. sativa* genome was described by Kagale et al. (2014) and is further specified herein. Subgenomes G1 (*n* = 6, N6) and G2 (*n* = 7, N7) mirror the genome structure of *C. neglecta* including the highly reshuffled chromosome Cn2 (Cs2 and Cs8, blocks M-Nb, Jb, Db, Ea, I, Eb, Ja, and Da; Figures 3 and 4). As the chromosome Cn2 originated through complex chromosome shattering, its repeated origins in the two *C. sativa* subgenomes are highly improbable. Thus, the presence of two Cn2 chromosomes in subgenomes G1 and G2 represents the strongest support for *C. neglecta*-like genome being the progenitor of both subgenomes (Figure 4; Supplemental Figure 8; Supplemental Data Set 1G). *C. neglecta*-specific fusion chromosome Cn6 (CAM6 + CAM7; blocks U, T, S, V, W, and X) was identified in

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**Figure 1.** (continued).

(B) to (D) Chromosomes of *C. hispida* (see [B]; Ch1 to Ch7), *C. neglecta* (see [C]; Cn1 to Cn6), and *C. laxa* (see [D]; C11 to C16) revealed by comparative painting using Arabidopsis BAC contigs as painting probes on pachytene and mitotic chromosome spreads. Chromosomes were counterstained with DAPI. The fluorescence of painting probes was captured as grey scale photographs and pseudocolored to match the eight AK chromosomes. Arrowheads point to centromeres. Bars = 10 μm.
subgenome G1 (chromosome Cs6), whereas ancestral chromosomes CAM6 and CAM7 are homeologous to Cs12 and Cs13 in subgenome G2 (Figure 4; Supplemental Figure 8). This fact, together with the number of chromosomes in the two subgenomes (n = 6 and n = 7 in subgenome G1 and G2, respectively), argues for an auto-allotetraploid C. neglecta–related parental genome formed by hybridization between two (sibling) taxa of C. neglecta distinguished by chromosome numbers (n = 6, N6 and n = 7, N7). Except for an ~10.67-Mb paracentric inversion on chromosome Cs11 [breakpoints within blocks O, between BACs F5I10 (AF195115) and F6N23 (AF058919), and R, T30N20 (AL365234)], subgenomes G1 and G2 did not experience subgenome-specific
Figure 3. Comparison of the Two Shattered Chromosomes in Camelina Diploids.

(A) Three-way comparison of GBs on chromosomes 2 and 4 in C. hispida (Ch, chromosomes Ch2 and Ch4), C. neglecta (Cn, Cn2, and Cn4), and C. laxa (Cl, Cl2 and Cl4). Color coding and capital letters correspond to three AK chromosomes (AK2, red; AK4, violet; AK5, orange) and six GBs (D, E, I, J, K-L, and M-N), respectively.

(B) Straightened pachytene chromosomes of C. hispida, C. neglecta, and C. laxa painted using differentially labeled Arabidopsis BAC contigs corresponding to six complete GBs. The fluorescence of painting probes was pseudocolored to match the corresponding AK chromosomes.
Figure 4. Genome Structure of *C. sativa*.

(A) Circos graphic displaying chromosomal collinearity between 20 chromosomes (Cs1 to Cs20) of the three *C. sativa* subgenomes (N6N7H, i.e., G1-G2-G3, left half of the circle) and 2 × 6 chromosomes of *C. neglecta* (Cn1 to Cn6) and seven chromosomes of *C. hispida* (Ch1 to Ch7, right half of the circle). Color coding and capital letters correspond to eight chromosomes and 22 GBs of ACK, respectively. See Supplemental Tables 2, 3, and 7 for boundaries of all GBs.
rearrangements and remained stable. As subgenome G3 (n = 7, H) resembles genome of C. hispida, including C. hispida–specific chromosomes Ch2 (Cs15), Ch4 (Cs17), Ch5 (Cs18), and Ch6 (Cs19), its structure provides conclusive evidence for the parentage of C. hispida. A single centromere repositioning on Cs16 (the Ch3 centromere, originally located between blocks G and H, was shifted ~1.54 Mb into block H, i.e., between BACs F16F14 (AC007047) and T2421 (AC005825)) differentiates the H subgenome from the parental C. hispida genome (Figure 4; Supplemental Figure 8; Supplemental Data Set 1G).

Altogether, our data suggest that the hexaploid genome of C. sativa (n = 20, N5N5H) originated through hybridization between the auto-allotetraploid C. neglecta (N5N5, n = 13) and diploid C. hispida (H, n = 7); the odd chromosome number in the inferred tetraploid C. neglecta was formed by hybridization between the ancestral (n = 7, N5) and derived C. neglecta–like genome (n = 6, N6), respectively.

To detect differences between hexaploid genomes of the cultivated C. sativa and its wild relative C. microcarpa, painting probes following the structure of 20 C. sativa chromosomes were identified in C. microcarpa. All 20 chromosome pairs were perfectly collinear in both hexaploid species (Supplemental Figure 8; Supplemental Data Set 1G). The identical chromosome number, subgenome composition, and chromosome structure indicate a common origin and recent genetic divergence of C. microcarpa and C. sativa.

Phylogenetic Analysis Further Supports the Allopolyploid Origin and Parentage of the Camelina Polyploids

To elucidate the phylogenetic relationships among the Camelina species, we provide additional support of parentage for polyploids and test possible hybridization events with 48 single-copy nuclear genes leveraged for the reconstruction of species trees and generation of phylogenetic networks. Four sets of data, with selective inclusion/exclusion of polyploid species, were analyzed (Supplemental Table 1): set 1 included all tested Camelina species, whereas the remaining data sets contained the diploids and (1) C. rumelica (set 2), (2) C. rumelica and C. sativa (set 3), and (3) C. rumelica and C. microcarpa (set 4).

Species trees were inferred for each set of data using the multi-species coalescent model in ASTRAL (Supplemental Figure 9) and Pseudo-likelihood for Estimating Species Trees (MP-EST; maximum pseudo-likelihood estimate implemented in Maximum species coalescent model in ASTRAL (Supplemental Figure 10). Two main clades were supported by the most probable phylogenetic networks for sets 1, 2, and 3 (Supplemental Figure 11).

Species trees based on set 4 differed between the two models. Whereas the ASTRAL tree was congruent with trees based on sets 1 and 3, the MP-EST tree retrieved C. hispida to be sister to C. rumelica, and C. laxa as sister to the remaining Camelina species. In both analyses, some clades had low statistical support (bootstrap value < 70%) due to reticulate evolution resulting from hybridization and polyploidization.

As the bifurcation branching pattern allowed in species trees cannot fully uncover complex evolutionary scenarios, we reconstructed phylogenetic networks in Phylogen testing for zero to three reticulation events. The most probable phylogenetic network, based on the all-species data (set 1) and assuming two hybridization events, is shown in Figure 5. These data suggest that C. laxa is the only species that did not participate in any hybridization event. The tetraploid C. rumelica originated through hybridization between diploid genomes closely related to C. hispida and C. neglecta. The second and independent hybridization between C. hispida (or closely related species) and C. neglecta led to the origin of a common hexaploid ancestor of C. sativa and C. microcarpa. The two independent hybridization events are also supported by the most probable phylogenetic networks for sets 3 and 4 (Supplemental Figure 11).

DISCUSSION

With detailed chromosomal analyses, we have characterized genomes of diploid Camelina species, identified genome-specific chromosomal signatures, and subsequently reconstructed the origin of Camelina allopolyploids. Our data corroborate the allohexaploid origin of C. sativa and C. microcarpa (Hutcheon et al., 2010) through a merger of three diploid (sub)genomes with seven or six chromosome pairs (Kagale et al., 2014).

By comparing genomes of the three extant diploid Camelina species (C. hispida, C. laxa, and C. neglecta), we have parsimoniously inferred the structure of an ancestral diploid Camelina genome (CAM, n = 7) preceding the divergence of diploid species. Although our CAM genome and the ancestral derivative of ACK genome, previously inferred from the C. sativa draft genome sequence (Kagale et al., 2014), are overall congruent, the CAM genome offers a higher precision of the ancestral genomic structures, including detailed characterization of chromosomal breakpoints. The CAM genome benefits from a more precise definition of ancestral GBs (Lysak et al., 2016) and incorporation of newly gained information on diploid genome structures. Furthermore, in derivative of ACK, descending dysploidy from n = 8 to n = 7 was interpreted as a fusion of chromosomes AK2 and AK4 (AK2/4) accompanied by several inversions. However, due to our analyses of diploid Camelina

Figure 4. (continued).

(B) GISH of gDNA of C. neglecta (red fluorescence) and C. hispida (green fluorescence) to 40 mitotic chromosomes in C. sativa. GISH revealed 26 chromosomes (N5N5 and N5N5) labeled by the C. neglecta probe, whereas 14 chromosomes were contributed by C. hispida (HH).

(C) CCP on pachytene chromosomes of C. sativa. Painting probes for AK7 and AK8 revealed the structure of the fusion chromosome Cs6. Note that due to the whole-genome triplication the used BAC contigs also labeled homoeologous regions on other Cs chromosomes. Differentially labeled painting probes are shown in experimental colors (red, green, and yellow fluorescence) and pseudocolored in turquoise and pink, respectively, following the color coding for AK chromosomes. Chromosomes were counterstained with DAPI. Bars = 10 μm.
Chromosome number. The tetraploid Camelina species, Camelina neglecta, was recognized only recently (Brock et al., 2019) and more effort is needed to establish the full distribution of this species, Camelina neglecta. A closely related, recently diverged, population or species differs from Camelina neglecta by a genome complement that is more similar to the genome complement of Camelina hispida. Both taxa are referred to as Camelina neglecta (as Camelina microcarpa) or Camelina hispida, both taxa are referred to as Camelina microcarpa (Douglas et al., 2015), Cardamine flexuosa (Douglas et al., 2015), Cucurbita microcarpa (N=5,12 is its exclusive chromosome number. The tetraploid Camelina rumelica (n=13, N^3H) originated through an interspecific hybridization between diploid genomes closely related to Camelina neglecta (n=6, N^6) and Camelina hispida (n=7, H). Comparison of both Camelina subgenomes with the respective parental genomes revealed much more extensive shuffling of the recessive N subgenome and overall conservancy of the dominant H subgenome; post-polyplid genome reshuffling was restricted only to the N-subgenome chromosomes, whereas a single paracentric inversion was detected in the H subgenome.

The hexaploid genome of Camelina sativa (as Camelina microcarpa is genomically identical with Camelina sativa, both taxa are referred to as Camelina sativa hereafter) originated in two hybridization steps (Figure 6). The initial genome merger was either an intraspecific hybridization between two Camelina neglecta plants with deviating chromosome numbers (n=7 and n=6) or hybridization between closely related, recently diverged, populations or species differentiated by a n-1 descending dysploidy (N^7, n=7 and N^6, n=6). The tetraploid species has hybridized with the diploid Camelina hispida to form the allohexaploid progenitor genome of Camelina sativa and Camelina microcarpa (N^6N^7H, n=20). In contrast to the Camelina rumelica genome, the two N and one H subgenomes in Camelina sativa remained remarkably stable since their merger: in the absence of any inter-subgenome rearrangements, only two intra-subgenome rearrangements differentiate subgenomes N^7 and H from the parental ones (a paracentric inversion on chromosome Cs11 and centromere repositioning on Cs16). As the N subgenome in Camelina rumelica was restructured, whereas two N subgenomes in Camelina sativa remained conserved, we conclude that the allotetraploid Camelina rumelica originated earlier than the N^6N^7 tetraploid and the allohexaploid genome of Camelina sativa.

The absence of inter-subgenome chromosomal rearrangements mediated by homeologous recombination in Camelina sativa, Camelina microcarpa, and even in the older Camelina rumelica genome (Figure 6; Kagale et al., 2014; this study) corroborates the lacking subgenome dominance and fractionation bias in Camelina sativa (Kagale et al., 2014). Thus, Camelina allopolyploids can be put on the list of allopolyploid plant genomes with a general subgenome stability, such as Arabidopsis suecica (Novikova et al., 2017), Capsella bursa-pastoris (Douglas et al., 2015), Cardamine flexuosa (Mandaková et al., 2014), Cucurbita species (Sun et al., 2017), or teff (Eragrostis teff; VanBuren et al., 2019). It was proposed that the subgenome differences in the composition, densities, and regulation of transposable elements (TEs; Freeling et al., 2012; Bird et al., 2018; Edger et al., 2019) might be the primary cause of biased subgenome fractionation. If parental genomes show a comparable diversity of TEs and/or have generally low proportion of TEs, it can be hypothesized that post-polyplid genome fractionation will be unbiased. Although this is attractive, more factors, such as homeologous exchanges causing aberrant meiotic pairing and reduced fertility (Gaeta and Pires, 2010), may play role in promoting long-term subgenome stability.

**Evolution of Camelina Genomes Was Accompanied by Chromosome Shattering**

Chromosome shattering, or chromothripsis, was recently discovered by cancer genome sequencing (Stephens et al., 2011). It is defined as an extensive intra-chromosomal cut-and-reassembly process accompanied by a high density of breakpoints and is known to be involved in both cancer and congenital diseases (Forment et al., 2012; Maher and Wilson, 2012; Korbel and Campbell, 2013). Multiple mechanisms of chromothripsis have been discussed, including evidence that abortive apoptosis, telomere erosion, mitotic errors, micronucleus formation, p53 inactivation, repetitive sequence, fragile site, or particular DNA
The CAM ($n = 7$) has evolved from an older genome with eight chromosomes ($n = 8$). The divergence of Camelina diploids was associated with species-specific genome reshuffling resulting in descending dysploidy in $C$. neglecta and $C$. laxa (both $n = 6$). Hybridization between two $C$. neglecta–like genomes with seven ($n = 7$, $N^7$) and six ($n = 6$, $N^6$) chromosomes led to the origin of an auto-allotetraploid $C$. neglecta–like genome ($n = 13$, $N^6N^7$). The allotetraploid
conformations may stimulate chromothripsis occurrence (Ivkov and Bunz, 2015; Fukami et al., 2017; Pellestor and Gatinois, 2019). The evidence that plants can undergo the same extreme chromosome shattering observed in some human cancers and developmental syndromes is rather scarce and not yet conclusive. Tan et al. (2015) showed that Arabidopsis embryos resulting from a cross between a mutant with weakened centromeres and a wild-type plant underwent chromothripsis causing a shattered chromosome. In an alloplasmic wheat line (Triticum aestivum), multiple recombination events between two nonhomologous chromosomes generated a shattered chromosome built of alternating segments of the two chromosomes (Zhang et al., 2008). In the present study, using fine-scale COP, we uncovered a complex, chromothripsis-like rearrangement as a key mechanism underlaying the formation of the shattered chromosome CAM2 in the ancestral CAM genome. The origin of CAM2 was associated with descending dysploidy from n = 8 in ACK to n = 7 in CAM preceding the species diversification in Camelina. Later, CAM2 underwent further independent species-specific chromosome shattering in C. laxa (Cl2) and C. neglecta (chromosome Cn2). Our data provide new evidence for chromothripsis-like events in naturally occurring plant genomes. As chromothripsis represents a mechanism of rapid and profound genome shattering, it might have the potential for rapid appearance of evolutionary novelties beneficial to their carriers. Conversely, the complex and abrupt nature of such catastrophic events may limit their offspring transmission and fixation. It can be hypothesized that some instances of chromosome shattering in plants occur in individuals (hybrids) heterozygous for gametocidal genes, that is, genes whose absence in gametes leads to chromosomal breakages and reduced gametic fitness. Although these chromosomal aberrations are frequently too severe, causing full gametic sterility, some semilethal rearrangements can become fixed after fertilization (Endo, 1990; Nasuda et al., 1998).

**Descending Dysploidies in Camelina Were Mediated by Unusually Complex Chromosomal Rearrangements**

The diploid Camelina genomes provide us with valuable insight into shuffling of an ancestral genome during species divergence. In Camelina diploids, descending dysploidy from n = 8 (ACK) to n = 7 (CAM), as well as dysploidy from n = 7 (CAM) to n = 6 (C. laxa), was mediated by complex chromosomal rearrangements, including chromothripsis-like events involving numerous breakages, inversions, shuffling of GBs, and a centromere loss. As CAM and all other ancestral genomes in lineage I/clade A have descended from ACK (Lysak et al., 2016), patterns of descending dysploidy can be compared across the clade. In contrast to Camelina, the reduction from n = 8 to n = 7 in Neslia paniculata (Camelineae; Lysak et al., 2006) was mediated by a single Robertsonian-like translocation, and a single end-to-end chromosome fusion in Boechera (Boechereae; Mandáková et al., 2015) and Descurainia (Descurainiae; Mandáková et al., 2017). Similarly, descending dysploidy from n = 8 to n = 6 in Hornungia alpina (Descurainiae; Lysak et al., 2006) was mediated by a single end-to-end chromosome fusion and nested chromosome insertion, respectively, and similar dysploidy in Smeleowskia altaica (Smelowskeae; Mandáková et al., 2017) and Turritis glabra (without tribal assignment; Lysak et al., 2006) were mediated by a single end-to-end chromosome fusion and Robertsonian-like translocation, respectively. Genomes of the two n = 6 Camelina species were formed through descending dysploidy with contrasting patterns. In C. neglecta, dysploidy from CAM was mediated by a single end-to-end chromosome fusion, whereas three fusion chromosomes in C. laxa originated by chromosome shattering. The number of breakpoints generated during chromothripsis-related dysploidies in CAM and C. laxa exceeds even the breakpoints during the most extreme descending dysploidy from n = 8 in ACK to n = 5 in Arabidopsis, mediated by a single end-to-end fusion and two Robertsonian-like translocations (Lysak et al., 2006, 2016).

**The Domesticated C. sativa Descended from the Weedy C. microcarpa**

Here, we show identical subgenome composition and chromosome structure in C. microcarpa and C. sativa. These findings further validate previous findings showing C. sativa as being domesticated from the wild hexaploid C. microcarpa (Brock et al., 2018). The identification of C. microcarpa as the pre-domesticate allows for further studies in the process of redomestication and introgression of traits via breeding. However, little is known about the process of domestication in this taxon, although future population genetic studies in C. microcarpa may elucidate the demographic history of C. sativa. Hybrid formation between C. microcarpa and C. sativa has been observed (Séguin-Swartz et al., 2013; Martin et al., 2018) and may prove to be a valuable tool in generating novel cultivars of C. sativa with drought tolerance, disease resistance, and earlier flowering.

**Camelina as a Model for Polyploidy and Evolutionary Studies**

While Camelina and Arabidopsis are closely related and belong to the same crucifer clade (lineage I or clade A), the mesohexaploid Brassiceae belongs to lineage II or clade B (Franzke et al., 2011). Thus, the univalved Arabidopsis genomic resources can be more easily exploited for studies of (post-)polyploidy genome evolution in Camelina than in Brassica. Moreover, due to the extinction of Brassiceae progenitor genomes and progressed rediploidization of Brassica genomes, these (sub)genomes cannot be compared with parental diploid genomes as it is feasible in the Camelina polyploid complex. Identifying diploid parental genomes of the

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Figure 6. (continued).

geneome of C. numelica (n = 13, N=H) was formed upon hybridization between genomes related to C. neglecta (n = 6, N=H) and C. hispida (n = 7, H). The allohexaploid progenitor genome of C. sativa and C. microcarpa (n = 20, N=2N=H) originated through hybridization between the auto-allotetraploid C. neglecta-like genome (n = 13, N=2N) and C. hispida (n = 7, H).
tetraploid and hexaploid Camelina species opens the door not only for studies of (post-)polyploid genome evolution but also for creating lines of C. sativa with improved traits and potential re-synthesis of polyploid Camelina genomes. With the ever-rising interest in false flax for aviation biofuels, production of high-value molecules, and as an omega-3-rich feedstock, the time is ripe to develop Camelina as a model system for the field of polyploidy research.

CONCLUSIONS

All three extant Camelina polyploids originated trough recurrent hybridization between paleogenomes identical or closely related to C. hispida (H genome) and C. neglecta (N genome). Whereas the allotetraploid C. rumelica genome shows signs of post-polyploid subgenome-specific fractionation, the three subgenomes in the allohexaploid C. sativa genome remained conserved since the genome merger. Remarkably, descending dysploidy preceding the origin of the ancestral CAM genome was accompanied by complex chromosome shattering involving numerous breakages, shuffling of GBs, and a centromere loss. Equally, the origins of fusion chromosomes in the C. laxa were mediated by chromosome shattering. The genome-shattering alterations, resembling those associated with human disorders, have been reported rarely in naturally occurring plant species. The identical genome structure of C. sativa and C. microcarpa points to their common origin and corroborates C. microcarpa as the wild hexaploid pre-domesticate of C. sativa. Identifying diploid parental genomes of the tetraploid and hexaploid Camelina species opens the door for creating lines of C. sativa with improved traits and potential re-synthesis of polyploid Camelina genomes and allows for adopting Camelina as a model system for polyploidy research.

METHODS

Plant Material

The list of investigated accessions and their origin is provided in Supplemental Table 2. The experimental plants were grown from seeds in a growth chamber under the following conditions: temperature 21\(^\circ\)C, daylength 16/8 h, light intensity 150 \(\mu\)mol/m\(^2\)/s. Young inflorescences were collected and fixed in freshly prepared fixative (ethanol:acetic acid, 3:1) overnight, transferred into 70% (v/v) ethanol, and stored at \(-20^\circ\)C until use. Young leaves were used directly or dried in silica gel. gDNA was extracted from harvested leaves of 22 Camelina accessions (Supplemental Table 2) using a NucleoSpin Plant II kit (Macherey-Nagel).

Chromosome Preparations

Chromosome spreads from young fixed flower buds, containing immature anthers, were prepared according to a published protocol (Mandáková and Lysak, 2016a). Briefly, selected flower buds were rinsed in distilled water (twice for 5 min) and citrate buffer (10 mM sodium citrate, pH 4.8; twice for 5 min), and digested in 0.3% (w/v) cellulose, cytohelicase, and pectolyase (all from Sigma-Aldrich) in citrate buffer at 37°C for 3 h. After digestion, individual anthers were dissected on a microscopic slide in 20 \(\mu\)L of 60% (v/v) acetic acid, spread on the slide, and placed on a metal hot plate (50°C) for \(-30\) s, and fixed in freshly prepared fixative (ethanol:acetic acid, 3:1). Suitable slides containing well-spread pachytene and/or mitotic chromosomes were post-fixed in freshly prepared 4% (v/v) formaldehyde in distilled water for 10 min and air dried. Preparations were treated with 100 mg/mL RNase (AppliChem) in 2 \(\times\) SSC at 37°C for 60 min, and 0.1 mg/mL pepsin (Sigma-Aldrich) in 0.01 M HCl at 37°C for 5 min, and then post-fixed in 4% (v/v) formaldehyde in 2 \(\times\) SSC for 10 min, washed in 2 \(\times\) SSC twice for 5 min, and dehydrated in an ethanol series (70, 90, and 100% (v/v), 2 min each).

DNA Probes

In total, 674 chromosome-specific BAC clones of Arabidopsis (Arabidopsis thaliana) grouped into contigs according to eight chromosomes and 22 GBs, were used. To determine fine-scale chromosome structures, uncover species-specific chromosome rearrangements, and precisely characterize breakpoints, after initial CCP experiments, some BAC contigs were split into shorter subcontigs. In the case of highly reshuffled chromosome regions, individual differentially labeled BAC clones were used. gDNAs of Camelina hispida, Camelina laxa, and Camelina neglecta were used as probes in GISH experiments in Camelina polyploids (Camelina microcarpa, Camelina rumelica, and Camelina sativa). Individual BAC clones and gDNAs were labeled with biotin-dUTP, digoxigenin-dUTP, or Cy3-dUTP by nick translation as described previously (Mandáková and Lysak, 2016b). In CCP experiments, the concentration of each labeled BAC clone DNA varied from 100 ng per BAC clone/slide in diploids (C. hispida, C. laxa, and C. neglecta), to 300 ng in tetraploid C. rumelica, and up to 500 ng in hexaploids (C. microcarpa and C. sativa). For GISH experiments, 150 ng of each gDNA per slide were used. Labeled BAC clones or gDNAs were pooled following a given experimental design, ethanol precipitated, dried using a desiccator, and dissolved in 20 \(\mu\)L of 50% (v/v) formamide and 10% (w/v) dextran sulfate in 2 \(\times\) SSC per slide.

Fluorescence in Situ Hybridization, Microscopy, and Image Processing

For CCP and fluorescence in situ hybridization, 20 \(\mu\)L of the hybridization mix was pipetted on a suitable slide and immediately denatured on a hot plate at 80°C for 2 min. For GISH, 20 \(\mu\)L of the probe was denatured in a microfuge tube at 90°C for 10 min, placed on ice for 10 min, pipetted on a slide, and denatured on a hot plate at 80°C for 2 min. Hybridization was performed in a moist chamber at 37°C for 32 to 48 h. Post-hybridization washing was performed in 20% formamide in 2 \(\times\) SSC at 42°C. The immunodetection of hapten-labeled probes was performed as described by Mandáková and Lysak (2016b) as follows: biotin–dUTP was detected by avidin–Texas Red (Vector Laboratories) and amplified by goat anti-avidin–biotin (Vector Laboratories) and avidin–Texas Red; digoxigenin-dUTP was detected by mouse anti-digoxigenin (Jackson ImmunoResearch) and goat anti-mouse–Alexa Fluor 488 (Invitrogen). Chromosomes were counterstained with 4',6-diamidino-2-phenylindole (DAPI, 2 \(\mu\)g/mL) in Vectashield. The preparations were photographed using an Axioimager M2 epifluorescence microscope (Zeiss) with a CoolCube camera (MetaSystems). Images were acquired separately for all four fluorochromes using appropriate excitation and emission filters (AHF Analysetechnik). The four monochromatic images were pseudocolored and merged using Photoshop CS (Adobe Systems) and cropped using ImageJ (National Institutes of Health) software. Pachytene chromosomes were straightened using the straighten-curved-objects plugin in ImageJ (Kocsis et al., 1991). Circular visualizations of chromosome-scale pseudomolecules were prepared using Circos (Krzywinski et al., 2009).

To identify chromosomes from individual subgenomes in Camelina polyploids, CCP was combined with GISH (CCP/GISH; Mandáková et al., 2014). After initial CCP experiment, hybridization patterns were analyzed and photographed, and painting probes were removed by washing the slides in 2 \(\times\) SSC and 20% (v/v) formamide in 2 \(\times\) SSC at 42°C (5 min each).
Subsequently, the slides were post-fixed in 4% (v/v) formaldehyde in 2× SSC for 10 min, washed in 2× SSC (twice for 5 min), and dehydrated in an ethanol series (70, 90, and 100% [v/v], 2 min each). The subsequent denaturation, hybridization and immunodetection steps were performed as described above.

Phylogenetic Analysis

In total, 48 nuclear markers (Supplemental Data Set 1C) targeting coding regions were selected based on Stockenhuber et al. (2015). Candidate markers were analyzed using BLAST searches against the genome sequence of *C. sativa* (PRJNA264159; Kagale et al., 2014) to identify conserved primer binding sites. The Access Array System (Fluidigm) was used to generate PCR amplicons for all the 48 markers, following the manufacturer’s protocol. PCR products were subsequently sequenced on the Illumina-MiSeq platform (2×300-bp paired-end reads). The quality check of raw reads was performed using FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Poor-quality reads and Illumina adapters were removed using Trimmomatic 0.36 (Bolger et al., 2014). The HybPiper 1.3.1 pipeline (Johnson et al., 2016) was used to extract haplotypes of homeologous sequences for each gene in all accessions analyzed. The set of python scripts performed three main steps: (1) reads were searched and sorted according to a target gene, (2) assembled contigs were aligned to the reference genome, and (3) potential paralogous sequences were identified. Four sets of data were generated (Supplemental Table 1): set 1 included haplotypes of all tested species and accessions, while sets 2 to 4 included consensus sequences for all diploid species and haplotype sequences for partial taxon sampling of *Camelina* polyploids; only *C. rumelica* in data set 2, *C. rumelica* and *C. sativa* in data set 3, and *C. rumelica* and *C. microcarpa* in data set 4. Arabidopsis, which is generally accepted to be paraphyletic to core *Camelina* species (Nikolov et al., 2019), was selected as the outgroup in the phylogenetic study.

Multiple alignments were generated for each marker using MAFFT v.7.407 (Katoh and Standley, 2013), using default settings, automatically trimmed using Phyutility 2.2 (Smith and Dunn, 2008) and manually checked in Geneious 11.1.5. ([https://www.geneious.com](https://www.geneious.com)). Maximum likelihood (ML) trees were inferred using RAxML 8.2.12 (Stamatakis, 2014) with GTR+G substitution model and rapid bootstrap with 100 replicates. The best ML gene trees were compared against 100 random trees using the Shimodaira–Hasegawa test implemented in RAxML to test sufficient phylogenetic signal in alignments, whereby only trees that passed the test were used in the subsequent step. RAxML gene trees are shown in Supplemental File. Low support branches (bootstrap < 10%) of gene trees were collapsed using the Newick utilities 1.6 (Junier and Zdobnov, 2010) to get higher accuracy of species tree analyses. The ASTRAL 5.6.3 (Rabiee et al., 2019) and MP-EST method implemented in the STRAW: Species Tree Analysis Web server (Shaw et al., 2013) were used to estimate species trees from a set of gene trees and bootstrap replicates. The branch support values of species trees were generated using 100 bootstrap replicates.

As a classical species tree cannot explain reticulate types of evolution, we analyzed potential hybridization scenarios in PhyloNet 3.6.9 (Wien et al., 2018). The maximum pseudo-likelihood approach was used, enabling testing of hybridization events under the influence of incomplete lineage sorting and is not as computationally exhaustive as ML method. The following settings were used: maximum number of reticulations (hybridizations), 0 to 3; number of optimal networks, 10; and gene trees bootstrap threshold, 50. The obtained phylogenetic networks were visualized and manipulated in Dendroscope 3.5.10 (Huson and Scornavacca, 2012).

**Supplemental Data**

**Supplemental Figure 1.** Structure of the ancestral *Camelina* genome (CAM, n = 7) and its origin from Ancestral Crucifer Karyotype (ACK, n = 8).

**Supplemental Figure 2.** Genome structure and the origin of *C. hispida* (n = 7) from CAM.

**Supplemental Figure 3.** Genome structure and the origin of *C. neglecta* (n = 6) from CAM.

**Supplemental Figure 4.** Genome structure and the origin of *C. laxa* (n = 6).

**Supplemental Figure 5.** Genome structure and the origin of *C. microcarpa*.

**Supplemental Figure 6.** Genome structure of *C. rumelica*.

**Supplemental Figure 7.** Genome structure and the origin of *C. rumelica*.

**Supplemental Figure 8.** Genome structure and the origin of *C. sativa* and *C. microcarpa*.

**Supplemental Figure 9.** Species trees of all data sets generated in ASTRAL.

**Supplemental Figure 10.** Species trees of all data sets generated in MP-EST.

**Supplemental Figure 11.** The most probable networks of all data sets generated in Phylonet.

**Supplemental Table 1.** Scheme of data sets used in the phylogenetic analyses.

**Supplemental Table 2.** Collection data of *Camelina* species used in the present study.

**Supplemental Table 3.** List of used markers with gene IDs and lengths of theoretical amplicons.

**Supplemental Data Set 1A.** Structure of the eight chromosomes and 22 genomic blocks (GBs, A-X) of Ancestral Crucifer Karyotype (ACK).

**Supplemental Data Set 1B.** Genome structure of *C. hispida*.

**Supplemental Data Set 1C.** Genome structure of *C. neglecta*.

**Supplemental Data Set 1D.** Genome structure of *C. laxa*.

**Supplemental Data Set 1E.** Genome structure of the inferred ancestral *Camelina* genome (CAM).

**Supplemental Data Set 1F.** Genome structure of *C. rumelica*.

**Supplemental Data Set 1G.** Genome structure of *C. sativa* and *C. microcarpa*.

**Supplemental File.** Maximum likelihood gene trees for all four sets.

**Accession Numbers**

Sequence data from this work can be found under the accession numbers listed in Supplemental Table 3.

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AUTHOR CONTRIBUTIONS
T.M., J.R.B., and M.A.L. conceived and designed the study. T.M. and M.P. performed the experiments. I.A.A. contributed to the interpretation of results. T.M., M.P., J.R.B., and M.A.L. wrote the article. All authors read and approved the final article.

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REFERENCES


