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The complex genome and adaptive evolution of polyploid Chinese pepper (*Zanthoxylum armatum* and *Zanthoxylum bungeanum*)

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Abstract

Zanthoxylum armatum and *Zanthoxylum bungeanum*, known as “Chinese pepper”, are distinguished by their extraordinary complex genomes, phenotypic innovation of adaptive evolution and species-special metabolites. Here, we report reference-grade genomes of *Z. armatum* and *Z. bungeanum*. Using high coverage sequence data and comprehensive assembly strategies, we derived 66 pseudo-chromosomes comprising 33 homologous phased groups of 2 subgenomes, including from autotetraploid *Z. armatum*. The genomic rearrangements and two whole-genome duplications creating large (~4.5 Gb) complex genomes with high ratio of repetitive sequences (>82%) and high chromosome number ($2n = 4x = 132$). Further analysis of the high-quality genomes shed lights on genomic basis of involutional reproduction, allomones biosynthesis and adaptive evolution in Chinese pepper, revealing a high consistent relationship between genomic evolution, environmental factors, and phenotypic innovation. Our study provides genomic resources and new insights for investigating diversification and phenotypic innovation in Chinese pepper, with broader implications for protection of plants under severe environmental changes.

Keywords: Chinese pepper (*Zanthoxylum armatum* and *Zanthoxylum bungeanum*), Polyploid, Identified subgenomes, Phenotypic innovation, Adaptive evolution

Introduction

The mechanisms underlying phenotypic innovation, diversification and adaptive evolution are key issues in plant biology. Differentiation of floral organs, biosynthesis of kairomones/synomones in pollen and co-evolution with insect pollinators, as well as functional diversification and neofunctionalization accompanying genomic evolution are considered core phenotypic innovations responsible for the rapid divergence of angiosperms (Mandel, 2019; Zhang et al., 2017a; Zhang et al., 2020a; Zhang et al., 2020c). Beside the evolution of reproduction, genomic evolution also provides a large amount of raw genetic materials for adaptive evolution in angiosperms during the different timing of global environmental changes (Wu et al., 2020). The warm, equable climate of the later Cretaceous and early Cenozoic was suited to rapid diversification, facilitated by the reproductive and growth advantages of angiosperms, with a concomitant diminution of the prominence of gymnosperms (Biffin et al., 2012; De La Torre et al., 2017; Guo et al., 2020). Recent research on macroevolutionary patterns in gymnosperms uncovered a resurgence of gymnosperm diversification and expansion in the late Cenozoic, driven by environmental heterogeneity, particularly in cooler and arid climatic conditions (Stull et al., 2021). The ecological factors in different geological eras might drive the emergence of different morphologies and evolutionary radiations more broadly in plants.

The *Z. armatum* and *Z. bungeanum*, often called the ‘Chinese pepper’, are the most famous spice in China, for the characteristics of ‘affinal medicine and diet’. The description of Chinese pepper can be traced back to *Book of Songs (Shi jing)* about 3000 years ago (Waley and Arthur, 1996). Over 30 classical prescriptions containing Chinese pepper were recorded in the TCM (traditional Chinese medicine) monograph (Zhang et al., 2017b). Chinese pepper has now become the largest planted woody spice crop globally, with a plantation of more than one million hectares in China. Though it belongs to genus *Zanthoxylum* (family Rutaceae), related to Citrus, Chinese pepper shares imperfect flowers, autonomous apomixis, and the production of diverse allomones (Fei et al., 2021a; Wang et al., 2021). The species-specific alkylamides creating a unique tingling sensation by activating two members of the transient receptor potential (TRP) channels (The Nobel Prize in Physiology or Medicine, 2021), TRPV1 and TRPA1, not hot as red pepper (*capsicum spp.*), or pungent as black pepper (*piper nigrum*), making them a novelty status in anti-herbivores evolution (Caterina et

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al., 1997; McNamara et al., 2005; Menozzi-Smarrito et al., 2009). These properties served as phenotypic limitation for molecular speciation and co-evolution with pollinators in Chinese pepper, which distinguished from the advantages of reproduction and diversification in early angiosperms evolution. Comparative genomic analysis has revealed that diploid *Z. armatum* experienced a whole-genome duplication (WGD) event after divergence from *Citrus* around 26.6 million years ago, as well as extensive expansion of genes related to arid adaption (Wang et al., 2021). The geographic distribution and evolution of *Z. bungeanum* is hypothesized to have been shaped by climatic oscillations during the Pleistocene (Feng et al., 2020). *Z. bungeanum* is widely distributed in subtropical and temperate regions, while *Z. armatum* is confined to subtropical frost-free regions in southwest China where the average annual temperature is 17 °C or higher (Fei et al., 2021a). The difference in species distribution between *Z. armatum* and *Z. bungeanum* may have been shaped by adaptation to cold stress.

High quality genome sequences can provide extraordinary data for addressing major issues ranging from agriculture to ecosystems. However, the high heterozygosity and extremely variable chromosome numbers of *Zanthoxylum* species complicate genome assembly and sub-genome identification. Although the draft genomes of *Z. armatum* and *Z. bungeanum* have been published recently, due to its large size and extremely genomic characteristics, both the draft release consisted of many fragmentary contigs/scaffolds (the contigs N50 of *Z. armatum* and *Z. bungeanum* are 0.34 and 0.41M, respectively (Feng et al., 2021; Wang et al., 2021)), which may result in inaccuracies in genomic studies. Besides, the subgenomes of allotetraploid *Z. bungeanum* also has not been identified. Here, we present high-quality genome assemblies of two Chinese pepper species (*Z. armatum* and *Z. bungeanum*). They represent the two typical species with: different type of extraordinary complex genomes (autotetraploid, allotetraploid), widely planted but with different adaptability to ecological factors, and the most valuable spices in Rutaceae family. The high quality assembly and accurate subgenome identification allows us to explore the relationship of genomic evolution, phenotypic variations and adaptive evolution in Chinese pepper. These results show a novel evolutionary path for biotic and abiotic adaptation of Chinese pepper, which provided an ideal opportunity to understand evolutionary adaptation to ecological factors in the new geological age.

Results

Assembly of high-quality Chinese pepper genomes with comprehensive strategy

Z. armatum and *Z. bungeanum* (Supplementary Figure 1) represent two widely distributed Chinese pepper species and carry the same karyotype: $2n = 4x = 132$ (Figure 1a-1d, Supplementary Figure 2, 3 and Supplementary Note 1). The genome size of *Z. armatum* was estimated to be ~4.4 Gb with a repetitive content of 79.54% (Supplementary Figure 4a, 4b and Supplementary Table 2). A slightly larger genome of *Z. bungeanum* was estimated to be ~4.6 Gb with a repetitive content of 80.60% (Supplementary Figure 4c, 4d and Supplementary Table 2).

To obtain a high-quality chromosome-level reference genome, we combined 1.3 Tb of PacBio sequence ($>260\times$ genome coverage and N50 of 30 Kb), 530 Gb high-throughput chromosome conformation capture (Hi-C) ($\sim 106\times$) sequence and 620 Gb Illumina data (Supplementary Table 1, 3 and 4) with Canu (Koren et al., 2017) and LACHESIS (Burton et al., 2013) based strategies to overcome the challenges posed by high heterozygosity and large genome size (Figure 1e and Supplementary Figure 5). Especially, in the part of chromosome assembly, we propose an assembly method of global clustering and then local multiple iterative clustering (Supplementary Note 2). This yielded a monoploid assembly of 4.39 Gb with a contig N50 value of 3.31 Mb for *Z. armatum*, covering 97.7% of the estimated 4.4 Gb genome, and an assembly of 4.63 Gb with a contig N50 of 10.74 Mb for *Z. bungeanum*, covering 97.3% of the estimated 4.65 Gb genome. In addition, the alignment of Illumina short-read (98.14% and 99.29% of mapping rate) and benchmarking universal single-copy ortholog (BUSCO) (98.1% and 98.5% completeness) validated the high-quality assembly of the *Z. armatum* and *Z. bungeanum* genomes (Supplementary Table 5-8 and Supplementary Figure 14). Obviously, these assembly results of *Z. armatum* and *Z. bungeanum* have significantly improved both in contiguity and completeness compared with the two recently published incomplete draft genomes (Feng et al., 2021; Wang et al., 2021) (Table 1).

Interestingly, for the *Z. armatum* genome, plotted Hi-C linkage shows that the chromosome groups are clear cut with 66 chromosomes comprising 33 homologous groups with two allelic chromosomes in each (Supplementary Figure 8). In addition, syntenic analysis in *Z. armatum* also revealed highly consistent gene order in the two allelic chromosomes (Figure 1f and

Supplementary Figure. 9). Based on these data, we inferred that *Z. armatum* is an autotetraploid with the karyotype of $2n=4x=132$ (AAAA). Accordingly, based on the collinearity with a diploid genome (Wang et al., 2021) (Supplementary Figure 10), we further identified the *Z. armatum* genome into two allelic chromosomes genomes (each consisting of 33 chromosomes; Supplementary Figure 11-13), which contain 90.0% and 84.7% BUSCO completeness, respectively (Supplementary Figure 14 and Supplementary Table 6).

To investigate sequence divergence and evolutionary relationships, the syntenic between A subgenome (consisting of 33 chromosomes) of *Z. armatum* and *Z. bungeanum* (consisting of 66 chromosomes, Supplementary Figure 15) were also explored, revealing the distinct 1:2 syntenic relationship between chromosomes of A subgenome and *Z. bungeanum* (Figure 1h and Supplementary Figure 16), while incomplete collinearity between interchromosomal of *Z. bungeanum* (Supplementary Figure 17). In consideration of high level of heterozygosity, high assembly accuracy by Canu (Koren et al., 2017) and syntenic relationship between chromosomes in the *Z. bungeanum* genome, we inferred that *Z. bungeanum* have polyploid characteristics different from *Z. armatum*, belong to the allotetraploid with karyotype of $2n=4x=132$ (BBCC). Subsequently, based on the syntenic and genetic distance (Zhang et al., 2021) between chromosomes of A subgenome in *Z. armatum* and homologous chromosomes in *Z. bungeanum* (Figure 1e and 1h, Supplementary Figure 16 and 18), we further phase the monoploid genome into B and C subgenomes (Figure 1g, h, Supplementary Figure 19-22), with N50 58.9 Mb of 1.96 Gb (96.5% BUSCO completeness) and N50 92.3 Mb of 2.48 Gb (92.4% BUSCO completeness), respectively (Supplementary Figure 14).

A comprehensive gene model prediction that integrated homology-based prediction, RNA-sequencing-assisted prediction and ab initio prediction for *Z. armatum* and *Z. bungeanum* identified a total of 65,195 (32,942 and 28,618 in A₁ and A₂ subgenomes) and 68,202 (31,074 and 28,596 in B and C subgenomes) protein-coding genes with 98.2% (87.0% and 81.7% in A₁ and A₂ subgenomes) and 98.7% (93.7% and 90.8% in B and C subgenomes) complete BUSCO genes as a whole, with more than 98.57% and 98.05% genes functionally annotated via searches of NR, GO, KEGG, Swiss-Prot and TrEMBL databases, respectively (Supplementary Note 3, Supplementary Figure 23, Supplementary Table 12 and 13). In addition, the annotated gene models showed high consistent with *A. thaliana* and other

Rutaceae species (Supplementary Figure 24 and Supplementary Table 11), indicating highly credible gene model inferences.

Phylogenetic position and genome size evolution of Chinese pepper

We next inferred the phylogenetic position and divergence times between *Z. armatum* and *Z. bungeanum* with all nine sequenced species in the genus *Citrus*, and one species from genus *Atalantia*, one from genus *Arabidopsis*, one from genus *Piper*, one representative orthologue from monocots and ANA-grade angiosperms. Here we selected the A and B, C subgenomes to represent *Z. armatum* and *Z. bungeanum* in this analysis. In total, 317 single copy genes were identified and used to construct phylogenetic relationships, via concatenated and multispecies coalescent approaches. The results indicate that the common ancestor of the A and B clade was phylogenetically a sister to the C subgenome, and the divergence time for A and B was estimated to be ~5.9 MYA (2.7–14.4 MYA), well after the allotetraploid formation of *Z. bungeanum* ~7.4 MYA (4.3–17.8 MYA) (Figure 2a and Supplementary Note 4).

The subgenome size of the *Z. armatum* and *Z. bungeanum* genome (assembled A = 2.2 Gb, B = 1.96 Gb and C = 2.48 Gb) is nearly 6-fold greater than that of the *Citrus* genome (average 357.6 Mb) (Figure 2b and Supplementary Table 14). Genome expansion in plants is driven by two major phenomena leading to sharp increases: polyploidization (whole genome replication, WGD) and proliferation of transposable elements (TE). The *Z. armatum* and *Z. bungeanum* genome assemblies were thus a good opportunity to study the drivers of genome expansion in *Zanthoxylum*. Firstly, we analyzed the synonymous/nonsynonymous substitution and WGD events in these A, B and C subgenomes. Interestingly, there may be a similar evolutionary rate among the subgenomes of *Z. armatum*. However, the subgenome B and C of *Z. bungeanum* shown an asymmetric evolution pattern (Figure 2c). In addition, distributions of synonymous substitutions per synonymous site (K_s) within genes in syntenic blocks clearly indicated that two WGD events occurred (Figure 2d and Supplementary Figure 25-27).

What's more, peaks at similar K_s values were identified in other Rutaceae species (Supplementary Note 5.1 and Supplementary Figure 28, 29), which suggests an ancient single WGD event probably shared among Rutaceae members and another is *Zanthoxylum* species-specific. Considering that *citrus* species have no recent WGDs except the shared ancient triplication by all core eudicots (WGT- γ) (Xu et al., 2013), we determined that a peak of 1.51

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in K_s distribution corresponds to the WGT- γ event. Given the mean K_s value (0.09) of A-C subgenomes and their divergence date T (7.4 MYA), we estimated the synonymous substitutions per site per year as $6.08\text{E-}9$ for *Zanthoxylum*, which dated the WGT- γ at around 123.4 MYA and the *Zanthoxylum* species-specific WGD at about 20.6 MYA (Figure 2d and Supplementary Note 5.2). Analysis of the WGT- γ and *Zanthoxylum* species-specific WGD genes (Supplementary Table 15) revealed those genes were enriched in various Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways (for instance, flavonoid biosynthesis, terpenoid backbone biosynthesis, MAPK signaling pathway, plant hormone signal transduction and plant-pathogen interaction pathway; Supplementary Note 5.3 and Supplementary Figure 30), suggesting their contribution to species divergence. The enrichment of the same KEGG categories in retained duplicates from two WGDs suggests that potential functional innovations associated with stress resistance might have benefited Chinese pepper at multiple times during evolution. High proportions of recent paralogs and *Zanthoxylum*-specific genes are all indicative of more frequent gene gain or expansion in *Z. armatum* and *Z. bungeanum*. The appearance of these paralogs at that time is intriguing and could be related to genome reorganization associated with TE expansion and/or removal.

Based on the chromosome-level assemblies, a total of 82.4% and 83.94% repetitive sequences were identified in *Z. armatum* and *Z. bungeanum* genomes, respectively. Among these repeats, 82.3% are classified as interspersed repeats and the predominant type of TEs was long terminal repeat (LTR) retrotransposons, accounting for 71.2% of the genome, including 15.93% LTR/*Gypsy* and 23.2% LTR/*Copia* retro-elements in *Z. armatum* (Supplementary Table 9). For *Z. bungeanum*, LTR retrotransposons account for 70.6% of the genome, including 16.2% LTR/*Gypsy* and 21.0% LTR/*Copia* (Supplementary Table 10). These TEs on both genome exhibit an apparently random distribution on the chromosomes, and an inverse correlation with gene density (Figure 1f, g, Figure 3a and Supplementary Figure 31, 32).

As much as 71.2% of the *Z. armatum* genome and 70.6% of the *Z. bungeanum* genome are composed of LTRs. The massive increase of *Ty1-copia*, and to a lesser extent *Ty3-gypsy*, LTR-retrotransposons accounts for most of the genome size differences among Chinese pepper and *Citrus*, *Atalantia* or *Fortunella* (Figure 2a). In addition, TEs are highly enriched

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in different genic regions of the *Z. armatum* compared with the corresponding regions of *Z. bungeanum* (Figure 3b). The distributions of insertion times showed that LTR retrotransposons in Chinese pepper have experienced continuing and more recent amplification bursts from 0–5 MYA (Figure 3c). LTR retrotransposons in the Chinese pepper were further sub-classified into lineages, of which *Ty1-copia Angela* and *Ty3-gypsy Athila* elements are being their major lineage, and *Ty1-copia Alesia* and *Ty3-gypsy Galadriel* elements are the least abundant (Figure 3d). Investigation of TE representation in Chinese pepper and *Citrus* subspecies confirmed that TE dynamics have shaped *Zanthoxylum* diversity through successive expansions and deletions (Figure 2a). To determine the historical dynamics of the different *Ty1-copia* and *Ty3-gypsy* retroelements in the *Zanthoxylum* genome, we analyzed the divergence of the reverse transcriptase (RT) and integrase (INT) sequences of different TE lineages, revealing different evolutionary patterns among lineages (Figure 3d, e). For example, *Angela* and *Athila* elements are all relatively young in *Z. bungeanum* than *Z. armatum* genome, consistent with either an intense and recent burst of insertion or a strong selection against *Angela* elements in *Z. bungeanum* (Figure 3e and Supplementary Figure 33).

We hypothesized that if TEs are related to phenotypic differences between species, their distribution should differ between *Z. armatum* and *Z. bungeanum*. To test this, we compared the distribution of TEs between *Z. armatum* and *Z. bungeanum*. TEs are highly enriched in different genic regions, especially in the promoter regions of genes (accounting for 37.0%, 38.1% and 39.1% of the total number of A, B and C subgenomes, respectively; Figure 3b), indicating that TEs could potentially contribute to the diversification of gene expression. To assess whether TEs are polymorphic in *Z. armatum* and *Z. bungeanum*, 25 *Z. armatum* accessions and 87 *Z. bungeanum* accessions (Feng et al., 2020) (Supplementary Table 16) were also scanned, with *Z. armatum* and *Z. bungeanum* as a reference genomes, respectively. We identified 11,875 (11,625 insertion and 250 absence) and 35,479 (34,623 insertion and 856 absence) polymorphic TEs among *Z. armatum* and *Z. bungeanum* genomes, with the vast majority of polymorphic TEs show different frequencies of insertion across accessions (Figure 3f).

Genomic variation and paleohistory of modern *Zanthoxylum* genomes

Alignment of the genomes of *Z. armatum* and *Z. bungeanum* were made to determine genomic divergence between the two representative accessions. A total of 47,882,821, 38,694,912, 40,146,029 single nucleotide polymorphisms (SNPs) and 9,778,819, 9,000,799, 9,460,872 small InDels were identified among the three (A, B, C) *Z. armatum* and *Z. bungeanum* subgenomes, respectively (Supplementary Table 17 and Supplementary Figure 34). The chromosomal distribution of SNPs and small InDels were throughout the genome, with no mutation hotspots detected (Supplementary Figure 35), but as expected most were concentrated in intergenic region (Supplementary Figure 36). These SNPs and InDels are inferred to possibly have functional effects on a total of 18,700 genes in subgenome A₁ of *Z. armatum* and 16,359 and 15,756 genes in subgenomes B and C of *Z. bungeanum*, and these genes are over-represented in several biological pathways including anthocyanin biosynthesis, fatty acid biosynthesis, flavonoid biosynthesis, biosynthesis of unsaturated fatty acids, diterpenoid biosynthesis and MAPK signaling pathway (Supplementary Figure 37).

The high-quality reference genomes allowed us to identify presence/absence variations (PAVs) by a direct comparative genome analysis of the two accessions. A mass of fusion/fission or inversion events involving *Z. armatum* and its homologs in subgenome of *Z. bungeanum* occurred and covered different chromosomal regions (Figure 1f-1h). In addition, we identified 191,750 and 182,663 segments in *Z. armatum* with total lengths of 1.02 Gb (17,867 genes) and 1.15 Gb (18,343 genes) that are absent in B and C subgenomes of *Z. bungeanum*, and 175,099 and 211,211 segments in B and C subgenomes of *Z. bungeanum* with total lengths of 1.0 Gb (17,035 genes) and 1.42 Gb (18,670 genes) that are absent in *Z. armatum*. GO and KEGG functional enrichment analyses revealed that most of the PAV genes are significantly highly enriched in flavonoid biosynthesis and terpenoid backbone biosynthesis (Fisher's exact test, false discovery rate [FDR] < 0.01) (Supplementary Figure 38). Interestingly, the pathways enriched by PAV genes between subgenomes are highly similar, but have different enrichment degrees. In addition, the PAV genes that are related to flavonoid biosynthesis and terpenoid backbone biosynthesis were tended to be expressed in fruits (Supplementary Figure 39).

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To assess the palaeohistory of modern *Zanthoxylum* genomes with high number of chromosomes, we collected chromosome information from six species, including *Citrus grandis*, *Citrus sinensis*, *Citrus unshiu*, *Poncirus trifoliata*, *Z. armatum* (A subgenome) and *Z. bungeanum* (B and C subgenomes), representing major lineages of the Rutaceae, and performed a comparative genomic investigation. Here, we firstly compared the genomes of above six species to ancestral eudicot karyotype (AEK), which used in opium poppy genome (Guo et al., 2018). The synteny dot plot (Supplementary Figure 40) and evolutionary scenario (Supplementary Figure 41) both illustrated that more syntenic copies of AEK segments in *Zanthoxylum* genomes than *Citrus*. Then, the Ancestral Rutaceae Karyotype (ARK), which takes into account gene conservation among above six species was structured into 30 protochromosomes containing 12,936 protogenes (Figure 4). Compared to ARK, species in the genus *Citrus* likely needed at least 21 chromosomal fusions and multiple reversals/translocations to reach its current structure of nine chromosomes. By contrast, *Z. armatum* and *Z. bungeanum* experienced a much more complex evolutionary history involving an additional WGDs and post-WGD rearrangements that finally shaped its karyotype of 33 chromosomes in subgenome. At least one chromosomal fission, six reversals and two translocations were shared in *Z. armatum* and *Z. bungeanum* after *Zanthoxylum*-specific WGD, and then at least one chromosomal fission, 10 reversals and one translocation to form present-day karyotypes (Figure 4). Interestingly, both *Z. armatum* and *Z. bungeanum* have some chromosome that no any synteny segments with AEK or ARK, suggesting complex chromosome rearrangement in *Zanthoxylum* genomes and need for genome sequencing of more representative species to obtain a perfect evolutionary history of Rutaceae ancestral chromosomes (Supplementary Figure 40, 41 and Figure 4).

Evolution of imperfect flowers and apomixis

Morphogenesis of floral organ is considered a key phenotypic innovation for the subsequent rapid evolution and diversification of angiosperms, sculpted by the ABCDE model of MADS-type floral homeotic genes. Here, we identified 187 and 325 MADS-box genes in *Z. armatum* and *Z. bungeanum* respectively, ascribed to the subfamilies of the ABCDE model, including A function: *APETALA 1* (*API*); B function: *AP3* and *PISTILLATA* (*PI*); C function: *AGAMOUS* (*AG*); D function: *AGAMOUS-like* (*AGL*); and E function: *SEPALLATA 1* (*SEPI*) for interacting with ABC function proteins (Supplementary Table 18). Among these, the

function A and D MADS-box class were significant expanded, while *PI* genes reduced with only 3 and 4 were identified in *Z. armatum* and *Z. bungeanum* respectively (Figure 5a). We also found high expression of function A and D MADS-box genes in fruit, while function B gene (*PI*) were absent (Figure 5b). According to the classic quartet model of flower organ identity, *PI* genes were an indispensable member for petals (AP1-AP3-PI-SEP) and stamen (AP3-PI-AG-SEP) development. The differential expansion/constriction and expression of MADS-box genes were consistent with the imperfect flower organs of Chinese peppers, sharing a similar phenotype of class B mutants that have sepals rather than petals in the second whorl and carpels rather than stamens in the third whorl (Figure 5c)(Theißen, 2001; Theißen and Saedler, 2001). In addition, the expansion and expression of *AGL11* suggest a sporophytic apomixis in carpel, which had been reported in recent study (Figure 5c)(Fei et al., 2021b).

Biochemical innovation of anti-herbivorous

Terpene volatiles, served as attractants for pollinators or/and as defense compounds against herbivores and phytopathogens were largely determined by the transformation of terpenoids from allomones to kairomones/synomones(Zhang et al., 2020c). Here, a series of genes involved in terpenoid backbone biosynthesis were found to be expanded in Chinese peppers, including *1-deoxy-D-xylulose-5-phosphate synthase (DXS)*, *1-deoxy-D-xylulose 5-phosphate reductoisomerase (DXR)*, *2-C-methyl-D-erythritol 2,4-cyclodiphosphate synthase (ispF)*, *geranyl diphosphate synthase (GPS)*, *4-hydroxy-3-methylbut-2-enyl diphosphate reductase (HDR)*. Terpene synthases (TPSs) are responsible for the biosynthesis of the various terpene molecules (Figure 5a). A total of 88 and 65 TPSs were identified in *Z. armatum* and *Z. bungeanum* respectively, assigned to five previously recognized TPS subfamilies in angiosperms: TPS-a, TPS-b, TPS-c, TPS-e/f and TPS-g (Figure 5d and Supplementary Table 19)(Chen et al., 2011). The expansion of genes related to terpene biosynthesis have contributed to the production of different volatile compounds for attracting pollinators or generating chemical defenses (Figure 5e). We next analyzed the aroma substances in the fruits and leaves of Chinese peppers. A total of 78 substances were identified, the major components of which were terpenoids. Although the contents and constituents of terpenoids from different species and habitats are different, they were allomones instead of kairomones/synomones, such as Limonene, Myrcene, and Pinene etc. (Supplementary Table

20). These allomones exert multi-defensive activities, such as protection against pathogens and herbivorous insects, and have contributed to adaptive evolution (Herde et al., 2008; Kappers et al., 2005; Van Poecke et al., 2001).

Chinese peppers synthesize some unique alkylamides, especially hydroxy- α -sanshool (HAS) and its derivatives, which create a strong tingling sensation, are natural defenses against herbivores (Xiong et al., 1997; Yang, 2008). The alkylamides are alkyl or aryl amides, biosynthesized from branched-chain amine and unsaturated fatty acids in acyl transfer reactions (Buitimea-Cantúa et al., 2020; Rizhsky et al., 2016). The expansion of genes related to unsaturated fatty acid biosynthesis and amino acid decarboxylation (Figure 5a) provided necessary precursors for alkylamides biosynthesis, such as *3-ketoacyl-CoA synthase (KCS)*, *3-oxoacyl-[acyl-carrier-protein] synthase (KAS)*, *acetyl-CoA acetyltransferase (ACAT)*, *fatty acid desaturase (FAD)* and *amino acid decarboxylase (AADC)*. Notably, the significant expansion of *BAHD-acyltransferases (BAHD-AT)* (contains 66 and 52 members in *Z. armatum* and *Z. bungeanum*, respectively), suggests that biosynthesis of alkylamides was positively selected during adaptive evolution (Figure 5a) (Wang et al., 2021).

We further investigate the insect-resistance of Chinese peppers using leaves as materials, which both contain allomones and hydroxy- α -sanshool. The starved *Helicoverpa armigera* (Lepidoptera: Noctuidae) fed on different combinations of leaves for 24 hours and found that the leaves of *Z. armatum* and *Z. bungeanum* were intact, while the leaves of other species were almost completely eaten (Figure 5f). The resistance of Chinese peppers to insects may be related to their unique content of secondary metabolites.

Phenotypic innovation related to stress tolerance

Another notable feature shared between the *Z. armatum* and *Z. bungeanum* genomes is the significant expansion of plant disease resistance genes (*R* genes), including *leucine-rich repeat receptor-like serine/threonine-protein kinases (LRR-RLK)* and disease resistance proteins, *cysteine-rich receptor-like protein kinases (CRK)*. We identified 1,312 and 1,101 *R* genes in *Z. armatum* and *Z. bungeanum* respectively, accounting for nearly 2% (only 0.5% in *citrus*, for comparison) of the total gene count (Figure 6a). We also found marked expansions of genes related to abiotic stress responses, such as WRKY transcription factors (*WRKY-TFs*), NAC transcription factors (*NAC-TFs*), and Zinc finger CCCH domain-containing protein

(ZC3H). In addition to gene expansions shared between the two *Zanthoxylum* species, we also noted significant differences in gene families potentially related to flavonoid metabolism and differential adaptation to cold stress (Figure 6a). Two important structural genes in flavonoid metabolism, *Chalcone synthase* (*CHS*) and *UDP-dependent glycosyltransferases* (*UGTs*), were significantly expanded in *Z. bungeanum*. With respect to the classic CBF/DREB pathway involved in cold stress, two *DREB* families were constricted in the *Z. armatum* genome, but expanded in *Z. bungeanum*.

The difference of geographic distribution and phenotype between *Z. armatum* and *Z. bungeanum* provide ideal model to analysis the relationship between gene expansion and adaptive evolution. Compare with the geographic distribution of *Z. armatum* in subtropical frost-free regions with warmly and stable climates, *Z. bungeanum* was widely distributed in subtropical and temperate regions with more severe climates. Phenotypically, we identified 28 flavonoids with differential accumulation in the fruit of between *Z. armatum* and *Z. bungeanum* by using the widely-targeted metabolomics approach (Chen et al., 2013) with six representative cultivars of *Z. armatum* and *Z. bungeanum* in different cultivation regions (Supplementary Figure 42). The composition and content of flavonoids in *Z. bungeanum* was significantly superior to *Z. armatum*, especially the glycosylated flavones, such as cyanidin 3-o-glucoside, cyanidin 3-O-rutinoside, and delphinidin 3-O-glucoside (Figure 6c, d, e and Supplementary Table 21). 75 genes involved in flavonoid metabolism with differentially expressed between *Z. armatum* and *Z. bungeanum* were identified. Among them, 61 genes were consistently highly expressed in *Z. bungeanum* tissues including *CHSs*, *CHIs*, *UGTs* and genes in flavonoid branch pathway (Figure 6b). Additionally, we found that genes related to flavonoid biosynthesis shown asymmetric express pattern between different subgenomes. For example, the expression of *CHSs* and *CHIs* were biased towards C subgenome, while downstream genes *F3Hs*, *FLSs*, *DFRs* and *UGTs* were biased towards B subgenome. Interestingly, some of these genes were also affected by TE (Figure 6b). Flavonoids are indispensable for adaptation of plants to environmental stresses, such as cold stress and UV injury (An et al., 2020; Ilk et al., 2015; Schulz et al., 2015). It is tempting to speculate that genomic evolution distinctions in flavonoid metabolism are causally connected to the geographical distinctions between the two *Zanthoxylum* species, with *Z. armatum* restricted

to frost free regions of southwest China but *Z. bungeanum* being adapted to wider areas with severe climates.

Evolutionary synthesis: the phenotypic innovation related to adaptive evolution

In this report, the ancestral whole-genome triplication of the Eudicots (WGT- γ at about 123.4 MYA) and *Zanthoxylum* species-specific WGD (20.6 MYA) events were revealed during Chinese pepper evolution. This vast amount of time during the two WGD events was accompanied by extensive climatic variability, with cool, arid climates. To investigate the evolutionary feature in response to climates of Chinese peppers, we divided duplicate genes into four types (genes from WGT- γ , WGD, tandem duplication and segmental duplication) to analysis the possible relationships between genome evolution (WGDs and TE bursting) and environmental factors related to phenotypic innovation. Phylogenetic and syntenic analyses of the MADS-box genes indicate that WGT- γ were followed by constriction of *PI* (B function genes for petals and stamen), whereas the WGD event generate the *AGL11* (D function genes for apomixis). We also revealed that the WGDs caused expansion of key structural genes for terpenoid backbone biosynthesis, such as *DXS*, *DXR*, and *HDR*. Whereas tandem duplications were largely responsible for *TPS* gene family expansion. Phylogenetic analysis indicated *TPS* genes are divided into TPS-a, -b, -c, -e/f, and g subfamilies, resulting in the diversification of specialized terpene allomones. *BAHD-AT* genes were expanded through the WGT- γ , which connected the fatty acid and amino acid metabolism to synthesize species-special alkylamides. The innovation of imperfect flower, apomixis, biosynthesis of allomones and alkylamides together represent characteristic reproductive features of excluding pollinating insects and anti-herbivorous in Chinese pepper, rather than the high reproductive efficiency of co-evolution with pollinator in classic evolution of angiosperm. Additionally, a noticeable evolutionary feature of Chinese peppers genome is the significant expansion of stress resistance genes, including *R genes*, *NAC-TFs*, *WRKY-TFs* etc. These results together might suggest a transition from rapid reproduction and growth to high resistance to the adverse environment during the evolution of Chinese pepper. A speculative model relating paleoclimate to modern Chinese pepper genomes and their gene family content is shown in Figure 7, which shown a consistent relationship between genomic evolution, phenotypic innovation in response to severe environmental factors in Chinese pepper.

Discussion

Here, we assembled two reference-grade genomes of Chinese pepper (*Z. armatum* and *Z. bungeanum*) through high-depth PacBio sequencing and manual assembly strategy of "clustering while removing redundancy". Compared with two published data (Feng et al., 2021; Wang et al., 2021), we improve the continuity and accuracy of the genome by as much as 10 times (Table 1). These assemblies enabling for a new era of Chinese pepper basic research and improvement efforts. Polyploidy is a widespread phenomenon in flowering plants at some point in their evolutionary history (Gaeta et al., 2007). Based on the syntenic and genetic distance (Zhang et al., 2021) between homologous chromosomes, we identified the subgenomes of autopolyploid *Z. armatum* and allopolyploid *Z. bungeanum*, and compared the heterogeneity of the subgenomes in the evolution of Chinese pepper. The high-quality genomes and subgenomes enable us explore the species-specific WGD events, the burst of LTR retrotransposons and severe chromosome fissions and fusions that both resulted in the complex and large genome of modern Chinese pepper. Among the sequenced Rutaceae species, Chinese pepper have the largest genome, the largest number of chromosomes and the highest proportion of repeat sequences. These data complement the resources for the genomic and evolutionary research of Rutaceae species.

The phenotypic innovation of reproductive assurance and rapid growth were the key to successful origin and diversification of angiosperm from the middle to late Cretaceous period, which was known as the "Abominable Mystery" of Darwin (Buggs, 2017; Li et al., 2019; Project, 2013). During this geological period with climate optimum, the phenotypic innovation of angiosperm mainly includes morphogenesis of floral organs, biosynthesis of kairomones/synomones, and loss of *R* genes (Guo et al., 2020). However, in recent years a growing body of researches has revealed that angiosperm adaptive evolution was driven by prevalent polyploidizations and severe climates selection. The trait morphological innovations were associated with stress response (Wu et al., 2020; Zhang et al., 2020b). Here, the genomic evolution of Chinese pepper shows a degeneration of reproductive strategy, include the imperfect flower, autonomous apomixis and reduced attraction of pollinator. Instead, the abiotic and biotic stress adaptation had been activated during the evolution of Chinese pepper. With the severe environment in the later Cenozoic, Chinese pepper evolutionary strategy might have undergone a transition from rapid reproduction and growth

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to high resistance to the adverse environment, which is consistent with recent work that describes the resurgence of gymnosperm diversification and expansion in the late Cenozoic (Figure 7)(Stull et al., 2021). Additionally, the genetic difference in flavonoid metabolism and responses to severe climates between *Z. armatum* and *Z. bungeanum* provide an ideal example to study the relationship between genomic evolution, environmental factors, and phenotypic innovation, which known as “Neo - darwinism” in evolutionary theory(Hancock et al., 2021; Noble, 2015). First, the metabolic analysis revealed superiority of flavonoid accumulation in *Z. bungeanum* which suggested the enhanced adaptation to severe climates(An et al., 2020; Schulz et al., 2021; Schulz et al., 2015). The differential accumulated flavonoids were consistent with highly expression of genes related to flavonoid metabolism in *Z. bungeanum*. In addition, the comparison of the genome-wide transcriptional levels of subgenomes also revealed the biased expression between subgenomes. The asymmetric evolution of the two subgenomes exhibited advantages of allopolyploidy over the autopolyploidy in *Z. bungeanum* for functional differentiation and neofunctionalization of flavonoids biosynthesis. These results suggested a consistent relationship between genomic evolution, phenotypic innovation in response to environmental factors in Chinese pepper. The highly relationship between intrinsic genomic evolution, extrinsic environmental factors (abiotic and biotic stress), and phenotypic innovation in Chinese pepper provided novelty data for the research of plant adaptive evolution. Compared with gymnosperm, the prevalent WGDs and polymorphic TEs of angiosperm provides more genetic materials for adaptive evolution, which then transforms into evolutionary advantage consistently.

In conclusion, the high-quality genomes of Chinese pepper (*Z. armatum* and *Z. bungeanum*) serve as the microevolutionary hallmarks of potential adaptation to environmental factors in the later Cenozoic. Our study provides critical insight on adaptive evolution underlying diversification and phenotypic innovation in Chinese pepper, with important broader implications for protection and utilization of plants in the new geological era under severe environmental changes, relevant to current global environmental changes.

Methods

Plant materials

The highly homozygous *Z. armatum* and *Z. bungeanum* were cultivated in the Sichuan Province, China (102.627533 E, 29.370615 N). For whole-genome sequencing and assembly, we collected the fresh young leaves from one single plant for each species and immediately frozen in liquid nitrogen. Total RNA for each species was extracted from leaf, root and fruit (mixed with floral organ) tissues in same plant that was used for genome assembly.

Library construction and sequencing

Genomic DNA from *Z. armatum* and *Z. bungeanum* was extracted using a modified CTAB method. Three size-selected DNA libraries for each species were constructed and sequenced using a HiSeq 2500 instrument with 2×150 bp paired-end reads.

To enable an optimal assembly of the large and complex (high heterozygosity and repetitive DNA) reference genome, more than 1.3 Tb of long sequence ($>260\times$ genome coverage and N50 of 30 Kb) for *Z. armatum* and *Z. bungeanum* were generated from SMRT cells on PacBio Sequel II platforms.

For the Hi-C sequencing and scaffolding, the fresh young leaves were fixed with formaldehyde and lysed, and the cross-linked DNA was then digested with DpnII. Hi-C DNA recovery and subsequent DNA manipulations were performed following Hu et al.(Hu et al., 2019). Then three Hi-C library were sequenced on an Illumina NextSeq instrument with 2×150 bp reads. A total of 530 Gb and 498 Gb raw data were produced for *Z. armatum* and *Z. bungeanum*, respectively.

Karyotyping and genome size estimation

For *Z. armatum* and *Z. bungeanum* karyotyping, chromosome painting techniques followed Gerlach et al.(Gerlach and Bedbrook, 1979; Gerlach and Dyer, 1980) with adaptations performed for root tissues. The dispersed cells of chromosome spreads at meiotic stages were counterstained with DAPI (4',6-diamidino-2-phenylindole) and fluorescence *in situ* hybridization with telomere repeats Oligo-(TTTAGGG)₆ as probes to determine the chromosome number. In addition, we used 5SrDNA and 18SrDNA repeats as probes for

fluorescence *in situ* hybridization to determine chromosome ploidy. The chromosome number and ploidy were photographed under an Olympus BX63 fluorescence microscope.

The genomic size of *Z. armatum* and *Z. bungeanum* was estimated combine with the flow cytometer approach and k-mer frequency analysis. For flow analysis, young leaves of *Z. armatum*, *Z. bungeanum*, *Gossypium hirsutum* (reference standard, 2.4 Gb) and *Gossypium raimondii* (reference standard, 737.8 Mb) were cut using a razor blade and incubated in staining solution (Sysmex) at room temperature for 5 minutes. The relative DNA content of isolated nuclei was analyzed using a flow cytometer, Sysmex CyFlow Cube8 (Sysmex Partec GmbH, Goerlitz, Germany), while data were acquired and processed by BD FACS DIVA software (v.7.0). Approximately 620 Gb Illumina sequence data for *Z. armatum* and *Z. bungeanum* were used to calculate K-mer frequencies by kmerFreq (<https://github.com/fanagislab/kmerfreq>) and estimate the genome size (G) with formula: $G = \text{k-mer number} / \text{k-mer depth}$ (Supplementary Figure 4 and Supplementary Note 2).

De novo assembly and phasing into subgenomes

The genome was assembled into chromosome-level in a step-wise manner as summarized in Figure 1e and Supplementary Figure 5. Raw PacBio reads were first self-corrected using an error correction module embedded in Canu (v.2)(Koren et al., 2017). The high-quality PacBio sub-reads were then used for contig-level assembly by using Canu(Koren et al., 2017) with parameter correctedErrorRate 0.050 and minReadLength 5000. We then used corrected PacBio long reads to polish the draft assembly using Arrow (<https://github.com/PacificBiosciences/GenomicConsensus>).

For chromosome-level assembly, these clean data were trimmed to remove low-quality bases and Illumina adapter sequences using trim-galore (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/), and then checked with HiCUP(Wingett et al., 2015). The clean HiC read pairs were aligned to consensus assembly described above using BWA-MEM(Li, 2013) with a default parameter. Subsequently, the uniquely mapped data were retained to perform cluster, order and orient the contigs by using LACHESIS software(Burton et al., 2013). Parameters for running LACHESIS included: CLUSTER_MIN_RE_SITES, 225; CLUSTER_MAX_LINK_DENSITY, 2; ORDER_MIN_N_RES_IN_TRUN, 105; ORDER_MIN_N_RES_IN_SHREDS, 105. In

order to overcome the challenges posed to HiC assembly by high heterozygosity and repetition sequences, as well as the genomic characteristics of multiple chromosome numbers, we propose an assembly method of global clustering and then local multiple iterative clustering in step of chromosome assembly using Hi-C reads (Supplementary Note 2). Finally, for each chromosome cluster, we inspect and manually correct with Juicebox assembly tools (<https://github.com/aidenlab/Juicebox>, v2.7.8).

To further improve the accuracy of reference assembled contigs, two-step polishing strategies were performed: we first used PacBio long reads and carried out gap filling with PBJelly(English et al., 2012) and then used highly accurate Illumina paired-end reads to further correct the base errors with Pilon (v.1.20)(Walker et al., 2014).

To evaluate the quality of the genome assembly, the Illumina sequencing reads were mapped use Bowtie2 (v.2.3.5)(Langmead and Salzberg, 2012). To evaluate the completeness of genome assemblies, the 1,614 conserved protein models in the embryophyta_odb10 dataset were searched against both genomes by using the BUSCO (v.5.0) program(Manni et al., 2021) with the --long parameter.

Genome annotation: repetitive sequences, gene models and noncoding RNA

The annotation of repetitive DNA followed both homology-based prediction and de novo identification of repeats as previously described(Hu et al., 2019). In brief, TRF (v. 4.07b)(Benson, 1999) and MISA(Beier et al., 2017) were used to identify tandem repeats and simple sequence repeats (SSRs). Long terminal repeats (LTRs) were identified using LTR_retriever(Ou and Jiang, 2018) on the basis of the results of LTRharvest(Ellinghaus et al., 2008) and LTR_Finder(Ou and Jiang, 2019) with the suggested parameters described in the manual. RepeatMasker (v. 4.0.5)(Chen, 2004) were utilized to search for known transposons (a *de novo* repeat library of *Z. armatum* and *Z. bungeanum* that built by RepeatModeler) and Extensive de-novo TE Annotator (EDTA)(Ou et al., 2019) was used for comprehensive TE identification. All aforementioned results were combined and merged to generate a non-redundant list of repeat elements reside in the genome.

A comprehensive process that integrated homology-based prediction, RNA-sequencing-assisted prediction and ab initio prediction were used for gene model prediction using

genome that all repetitive regions have been soft-masked. For ab initio prediction, Augustus (v3.3.1) (Stanke and Waack, 2003), Genscan (v.3.1) (Burge and Karlin, 1997), GeneID (v.1.4) (Blanco et al., 2007), GlimmerHMM (v.1.2) (Majoros et al., 2004), GeneMarkS-T (v.4) (Besemer et al., 2001) and SNAP (v.2006-07-28) (Korf, 2004) were used for de novo-based gene prediction with the default parameters. Additionally, filtered proteins (incomplete and wrong) of five species (*A. thaliana*, *C. clementina*, *C. sinensis*, *C. unshiu* and *P. trifoliata*) were used for homology-based prediction with GeMoMa (v1.5.3) (Keilwagen et al., 2018) and GeneWise (v.2.4.0) (Birney et al., 2004) using the default settings. Then, PASA (v2.4.1) (Haas et al., 2008) was used for RNA-seq and Iso-Seq based gene prediction. Finally, the results from the three approaches were integrated using EvidenceModeler (EVM; v1.1.1) (Haas et al., 2008) to obtain the raw gene set. To obtain a precise gene set, some genes whose sequences included transposable elements were filtered with TransposonPSI software (<http://transposonpsi.sourceforge.net>). To assess the completeness of the gene set, BUSCO (v.5.0) (Manni et al., 2021) was used to evaluate the gene set based on the encoded proteins using *embryophyta_odb10*.

Putative gene functions were assigned using the best match to SwissProt and TrEMBL databases using BLASTP (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). Motifs and domains were searched using InterProScan (v.5.52) (Jones et al., 2014) against all default protein databases including ProDom, PRINTS, PfamA, SMART, TIGRFAM, PrositeProfiles, HAMAP, PrositePatterns, SITE, SignalP, TMHMM, Panther, Gene3d, Phobius, Coils and CDD. The gene pathways of the predicted sequences were extracted from the Kyoto Encyclopedia of Genes and Genomes (KEGG) Automatic Annotation Server (v2.1).

tRNA-encoding genes were predicted by tRNAscan-SE (v.1.3.1) (Lowe and Chan, 2016). MicroRNA and small nuclear RNA (snRNA) genes were found by searching against the Rfam database (release 12.0) with Infernal (v.1.1.1) (Nawrocki and Eddy, 2013).

Ancestral karyotype reconstruction

The ancestral karyotype of Rutaceae was reconstructed using *Citrus grandis*, *Citrus sinensis*, *Citrus unshiu*, *Poncirus trifoliata*, *Z. armatum* and *Z. bungeanum*, for which have complete and chromosome-level genome assemblies, as previously described (Murat et al., 2017). In brief, *Z. bungeanum* was adopted as a reference genome and BLAST (e-value <1e-5) was

used for two-by-two interspecies comparisons. The parameters cumulative identity percentage (CIP) and cumulative alignment length percentage (CALP)(Throude et al., 2009) were used to filter results of pairwise sequence alignments and to define conserved/duplicated gene pairs (putative protogenes, or pPGs). The pPGs conserved in all six genomes (core protogenes, or core pPGs) were extracted and these core pPGs are used to identify orthologs coordinate of synteny blocks (SBs) by MCScanX(Wang et al., 2012) with the filtering out of groups of fewer than five (pPGs) genes. Then GRIMM-Synteny(Tesler, 2002) was used to merge the coordinates into SBs correspondence between the six collinear groups and MGR (v.2.0.1)(Lin et al., 2009) was used to rearrange the SBs into ancestral protochromosomes (also referred to as contiguous ancestral regions (CARs)) on the basis of chromosome-to-chromosome orthologous relationships between the compared genomes.

Identification of SNPs, small indels and PAVs

The homologous pseudochromosome sequence between the *Z. armatum* and *Z. bungeanum* was aligned with MUMmer (v3.23). SNPs and small indels (length <100 bp) was identified using Show-SNPs. Both inversions and translocations were identified with a length of >100 bp. PAVs were extracted by scanPAV(Giordano et al., 2018) with default parameters and the resulting PAVs that shorter of 1,000 bp were filtered out as noise.

Phylogenetic analysis and divergence time estimation

The assembled genome of *Z. armatum* and *Z. bungeanum* allowed us to understand its evolution and to estimate divergence time within Rutaceae species. In order to achieve a robust phylogenetic reconstruction with high confidence and concordance, we used gene models from Rutaceae and other plant species, which include all nine sequenced species in the genus *Citrus* (*Citrus clementina*, *Citrus grandis*, *Citrus ichangensis*, *Citrus medica*, *Citrus reticulata*, *Citrus sinensis*, *Citrus unshiu*, *Fortunella hindsii* and *Poncirus trifoliata*), one from *Atalantia* (*Atalantia buxifolia*), one from *Arabidopsis* (*Arabidopsis thaliana*(The Arabidopsis Genome, 2000)), one from *Piper* (*Piper nigrum*(Hu et al., 2019)), one representative orthologue from monocots (*Oryza sativa*(Goff et al., 2002)) and ANA-grade angiosperms (*Nymphaea colorata*(Zhang et al., 2020a)), together with *Zanthoxylum armatum* and *Zanthoxylum bungeanum* to putative the orthologous gene groups. The longest transcript was selected to represent each gene. ORFs with premature stop codons, that were not

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multiples of three nucleotides long, or encoded less than 50 amino acids, were also removed. Then OrthoMCL(Li et al., 2003) was used to construct gene families based on all-against-all BLASTP alignment among the 16 species. The total of single-copy orthologous gene sets were extracted from the above gene family analysis and aligned using MAFFT (v.7.471)(Kazutaka and Standley, 2013), then low-quality regions were trimmed using Gblocks (v.0.91b)(Talavera and Castresana, 2007). A maximum likelihood phylogenetic tree was constructed using concatenated alignment with RAxML (v8.2.1264) and the PROTGAMMAILGF model to automatically determine the best reasonable tree by conducting 1,000 bootstrap replicates. The concatenated alignment and maximum likelihood tree that used as a starting tree were input into BEAST (Bayesian Evolutionary Analysis Sampling Trees2; v2.1.2) and MCMCtree program(Yang, 2007) to estimate species divergence time. A Calibrated Yule model with a Strict Clock rate and gamma hyperparameter of prior distribution were used to estimate the divergence time. Speciation event dates for *Citrus-Atalantia* (Normal model, Mean: 22.5 MYA, Sigma: 0.5)(Peng et al., 2020), monocots-eudicots (Log Normal model, Mean: 185 MYA, Std dev: 9MYA)(Zhang et al., 2020a) and Magnolids-monocots-eudicots-ANA-grade angiosperms (Log Normal model, Mean: 226 MYA, Std dev: 1MYA)(Zhang et al., 2020a), were used to calibrate the divergence time. The Markov chain Monte-Carlo analysis was repeated 10,000,000 times with 1,000 steps.

The expansion and shrinkage of gene numbers in different gene families between above species were estimated using CAFE (v. 4.2)(De Bie et al., 2006). A family-wise *p*-value (based on a Monte-Carlo re-sampling procedure) of 0.01 was used to indicate whether has a significant expansion or contraction occurred in each gene family across species.

Genome alignment and gene synteny analysis

Genome alignment either inter-species or intra-species of *Z. armatum* and *Z. bungeanum* was performed using the minimap2 (v.2.16-r922)(Li, 2018) program with parameters settings -x asm5 and dot plot was used to display the synteny block located in inter-chromosomal or intra-chromosomal.

To understand *Z. armatum* and *Z. bungeanum* genome evolution, synonymous substitutions per site (*K_s*) distribution of whole-genome duplication and segmental duplications in each

genome was investigated. First, the paranome was constructed by performing an all-against-all protein sequence similarity search using BLASTP with an E value cutoff of 1×10^{-10} , after which the reciprocal best hit (RBH) pairwise sequences of paralogous and orthologous relationships were identified and the protein sequences and corresponding codons sequences were aligned using ClustalW(Larkin et al., 2007) and PAL2NAL(Suyama et al., 2006), respectively. The K_s values were calculated using the YN model in KaKs_Calculator (v. 2.0)(Wang et al., 2010). Subsequently, K_s distribution was fitted using Gaussian mixture models (GMM) in the R package mclust (v.5.3) and used to examine the most recent WGD event in *Z. armatum* and *Z. bungeanum*. Secondly, we performed synteny analysis on *Z. armatum* and *Z. bungeanum* genes using MCScanX(Wang et al., 2012) with default parameters from top five self-BLASTp hits. The K_s for gene pairs located in syntenic block was calculated and K_s distribution was fitted and display using same method with above description.

The time of a WGD event was inferred by using the relative divergence of the duplicates and the formula $\text{divergence date} = K_s / (2 \times r)$, where r refers to K_s /year rate of *Zanthoxylum*.

Analysis of repeat and potential LTR bursts

Each of the whole genomes was searched for repeat annotation use EDTA and then refined the transposon-protein-domains using DANTE (RepeatExplorer server; <https://repeatexplorer-elixir.cerit-sc.cz/>). The hits were filtered to cover at least 80% of the reference sequence, minimum identity of 35% and minimum similarity of 45%, allowing for a maximum of three interruptions (frameshifts or stop codons). TE classes were defined according to Wicker et al.(Wicker et al., 2007) and TE lineages were defined according to Neumann et al.(Neumann et al., 2019). The polymorphic TEs of Chinese pepper were identified using TEMP2(Yu et al., 2021) based on the raw reads of 25 resequenced genomes of *Z. armatum* accessions(Feng et al., 2020) with the *Z. armatum* genome as the reference and 87 resequenced genomes of *Z. bungeanum* accessions(Feng et al., 2020) with the *Z. bungeanum* genome as the reference (Supplementary Table 16).

Full-length LTR-RTs were identified using EDTA and categorized into the subgroups of Copia-like and Gypsy-like were used to estimate the insertion time. The long terminal repeats for each LTR were extracted and aligned with MUSCLE (v.3.8.31), and the nucleotide

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distances (K) between them were calculated by the Kimura Two-Parameter approach using the distmat programme in EMBOSS (v.6.6.0). The insertion time (T) of each LTR was calculated by the formular: $T = K / (2 \times r)$, where r refers to a nucleotide substitution rate of per site per year in *Zanthoxylum* calculated by combination with species divergence time and K_s distribution.

The amino acid sequences of Copia-like and Gypsy-like superfamilies were aligned using MAFFT (v.7.471) (Kazutaka and Standley, 2013) with default parameters. The phylogenetic trees were constructed using fastTree (<http://www.microbesonline.org/fasttree/>).

RNA-seq library construction, sequencing and data normalization

Total RNA was extracted from leaf, root and fruit (mixed with floral organ) samples from *Z. armatum* and *Z. bungeanum*. After DNase treatment, RNA-seq libraries were constructed and sequenced on the Illumina HiSeq 2500 platform with 150 bp paired-end sequences according to the manufacturer's recommended protocol. A mix of total RNA was also selected to prepare a 20 Kb SMRTbell Template library and prepared for full-length transcriptome sequencing using the PacBio Iso-Seq protocol. The PacBio Iso-Seq3 pipeline (<https://github.com/PacificBiosciences/IsoSeq3>) was employed to obtain high-confidence transcriptome reads and used in gene annotation.

For paired-end raw reads, trim-galore was applied to remove adapters and poor-quality reads. The processed reads for *Z. armatum* and *Z. bungeanum* were aligned to its reference genome using HISAT2 (v.2.1.0), and quantification of gene expression (RPKM, TPM, and expression count data) was performed with StringTie (v.2.1.4).

Analysis of genes involved in the floral organ patterning

To identify putative MADS-box family genes in *Z. armatum* and *Z. bungeanum*, we searched the predicted proteome of *Z. armatum* and *Z. bungeanum* using hmmsearch in HMMER (v.3.0), based on the seed SRF-TF domains (PF00319) from the Pfam database. The Simple Modular Architecture Research Tool (SMART) and conserved domain databases were used to examine all candidate MADS-box genes and genes with incomplete MADS-box domains were removed. MADS-box classification was based on sequence similarity searches of identified MADS-box genes from *A. thaliana* (The Arabidopsis Genome, 2000) and *N.*

colorata(Zhang et al., 2020a). For evolutionary analysis of MADS-box genes, we aligned protein sequences using MAFFT (v.7.471)(Kazutaka and Standley, 2013) with E-INS-I iterative refinement method and automatically trimmed by trimAl (v.1.1)(Silla-Martínez et al., 2009). The alignment was then used to construct maximum likelihood phylogenetic tree using IQ-TREE (v.2.1.3)(Minh et al., 2020).

Floral scent measurement

In order to determine the aroma components and flavonoids metabolites contents of different Chinese pepper varieties, representative species from Chongqing, Shaanxi, Hubei, Yunnan and Gansu were selected (Supplementary Figure 42) to perform an LC-MS-based metabolomic analysis for the leaf according to the published report(Chen et al., 2013). In brief, samples from Chinese pepper were frozen in liquid nitrogen and ground to a powder. The volatile substances were extracted from 0.50 g powdered samples by Hydro-distillation for 3 h using a Clevenger type apparatus. GC-MS analysis was performed on an Agilent 7890B GC system with an Agilent Technologies 5977C Inert XL Mass Selective Detector, equipped with an HP-5MS UI column (30 m×0.25 mm×0.25 μm; Agilent Technologies). The essential oil components were identified by comparison of mass spectra with the NIST 2011 library data.

For LC-ESI-MS/MS analysis of flavonoids, 1 mL pre-cooled extractant (70% methanol aqueous solution) was added to powdered samples. The mixture was centrifuged at 12,000 r/min at 4 °C for 10 min. The supernatant was used for LC-MS/MS analysis. The extracts were analyzed using an LC-ESI-MS/MS system (UPLC, Shim-pack UFLC SHIMADZU CBM A system, <https://www.shimadzu.com/>; MS, QTRAP 6500+ System, <https://sciex.com/>). LIT and triple quadrupole (QQQ) scans were acquired on a triple quadrupole-linear ion trap mass spectrometer (QTRAP), QTRAP 6500+ LC-MS/MS System, equipped with an ESI Turbo Ion-Spray interface, operating in positive and negative ion mode. The ESI source operation parameters were as follows: source temperature 500 °C; ion spray voltage (IS) 5500 V (positive), -4500 V (negative); ion source gas I (GSI), gas II (GSII), curtain gas (CUR) was set at 55, 60, and 25.0 psi, respectively; the collision gas (CAD) was high. Instrument tuning and mass calibration were performed with 10 and 100 μmol/L polypropylene glycol solutions in QQQ and LIT modes, respectively. A specific set of MRM

transitions were monitored for each period according to the metabolites eluted within this period. Three biological replicates were performed in each experiment.

Data availability

The *Z. armatum* and *Z. bungeanum* assembly and annotation data are available at figshare (https://figshare.com/articles/dataset/Genome_Data_of_Chinese_pepper/20217635). The raw sequencing data used for *de novo* whole-genome assembly are available from the Sequence Read Archive under accession number PRJNA771757 and PRJNA771946. Transcriptome data of Illumina RNA-seq and PacBio Iso-Seq are available at the Sequence Read Archive under accession number PRJNA773368 and PRJNA773367. Further details on data accessibility are outlined in the supplementary materials and methods.

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Conflicts of interest

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

Author contributions

C.H., J.W., S.Z., P.X., H.Z., S.J., W.C. and Q.W. designed and supervised the research. Z.X., P.X. and M.M. performed the genome assemblies, annotation, transcriptome and phylogenetic analysis. F.W. and G.W. performed flow cytometer approach to estimate genome size. L.H., R.F., X.Q., L.Y., H.Z. and X.J. collected materials for sequencing and generated transcriptome data. Z.X., P.X. and L.H. analyzed the RNA-seq data. W.C. conducted the GC-MS analyses. J.W., H.D., K.L., S.J. and X.Z. provided constructive comments and suggestions on data analysis. Z.X. and L.H. wrote the manuscript with input

from all other authors. J.W., H.D., K.L., S.S. and S.J. edited the paper. All authors have read and approved the manuscript.

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FIGURE LEGENDS

Figure 1 **Major features of two Chinese pepper genome**. Discrimination of chromosomes within the *Z. armatum* (a) and *Z. bungeanum* (c) karyotype using FISH with telomere sequence (TTTAGGG)₆ as probe (green). Scale bar, 5 μ m. Three independent biological replicates. Ploidy-estimating of chromosomes within the *Z. armatum* (b) and *Z. bungeanum* (d) karyotype using FISH with 5SrDNA sequence as probe (red). Scale bar, 5 μ m. Three independent biological replicates. (e) Genome assembly and subgenomic identification strategy in Chinese pepper. Circos view of the *Z. armatum* (f) and *Z. bungeanum* (g) genome. Lanes depict circular representation of pseudomolecules (a) and the density of GC, repeat, LTR, genes and expression of fruit, root, leaf-specific genes (b–g). Lines in the inner circle represent links between synteny-selected paralogs. (h) Genome alignment of the B (Zb_B) and C (Zb_C) subgenomes in *Z. bungeanum* with A (Za_A) subgenome in *Z. armatum*. Lines

between chromosomes show syntenic regions and distinct 1:2 syntenic relationship between *Z. armatum* and *Z. bungeanum* were observed.

Figure 2 Rutaceae phylogenomics. **a**, Summary phylogeny and timescale of Rutaceae plant species. Blue bars at nodes represent 95% credibility intervals of the estimated dates. Evolution as expansion, gain, contraction or loss of the orthogene families in context of phylogenetic profile were located on the right of species tree. Barplots of the percentage (the total length divided by genome size) of TE and exon of the species corresponding to species tree were located on the far right. **b**, Number of gene-coding sequences (CDS) against genome size (Gb) for selected Rutaceae and outgroup sequenced genomes. Data points are represented by centered labeled boxes; overlapping points are indicated. **c**, Boxplot of the Ka/Ks ratio distribution of protein-coding genes in 66 chromosomes of *Z. armatum* and *Z. bungeanum*. The central line for each box plot indicated median. The top and bottom edges of the box indicated the 25th and 75th percentiles and the whiskers extend 1.5 times of the interquartile range beyond the edges of the box. **d**, Synonymous substitution rate (*Ks*) distribution plot for paralogs and orthologs of Chinese pepper with other outgroup species as shown through colored dotted and continuous lines, respectively. The arrow highlights the recent whole-genome duplication identified in Chinese pepper genome.

Figure 3 TE evolution in the Chinese pepper genome. **a**, Distribution of TE across the *Z. armatum* and *Z. bungeanum* genomes. **b**, Number of TEs in different genic regions of the *Z. armatum* and *Z. bungeanum* genomes. **c**, Analysis of intact LTR numbers and insertion time in *Z. armatum* and *Z. bungeanum* plants. **d**, Average age of TEs in *Z. armatum* and *Z. bungeanum* were revealed for the different lineages through RT and INT protein domains. The bar plot columns in right give the number of RT and INT domains present in the *Z. armatum* and *Z. bungeanum* genomes. **e**, Neighbor-joining (NJ) trees were built from RT domain sequence similarities among different lineage-specific copies identified in the *Z. armatum* and *Z. bungeanum* genome. Deep branching revealed ancient expansion while flat branching is consistent with a recent burst of insertion activity. Red branches correspond to outgroup sequences. **f**, Frequency distribution of polymorphic TEs (absence and insertion) in the genomes of 25 resequenced genomes of *Z. armatum* accessions with the *Z. armatum*

genome as the reference and 87 resequenced genomes of *Z. bungeanum* accessions with the *Z. bungeanum* genome as the reference.

Figure 4 Evolutionary history of chromosome in Chinese pepper. Evolutionary scenario for modern Rutaceae (*C. grandis*, *C. sinensis*, *C. unshiu*, *P. trifoliata*, *Z. armatum* and *Z. bungeanum*,) from the ARK of 30 protochromosomes. The modern genomes are illustrated at the bottom with the different colours reflecting the origin from the 30 ancestral chromosomes from the n=30 ARK (top). Duplication events are shown with a stars with different color, along with the chromosome reversal (R), translocation (T), fusions (Fu) and fissions (Fi) events. The time scale is shown on the left (million years).

Figure 5 Phenotypic innovation of floral morphogenesis and allomone biosynthesis in Chinese pepper. **a**, Expansion of genes involved in floral morphogenesis, terpene metabolism and alkylamide biosynthesis. **b**, Expression of MADS genes involved in floral morphogenesis. The gradient color for each gene represents the gene expression levels in root, leaf and fruit tissues of *Z. armatum* and *Z. bungeanum*. **c**, The ABCDE model of floral morphogenesis in Chinese pepper. **d**, Phylogeny of terpene synthases (TPSs) from Chinese pepper using a maximum likelihood tree. Branches are coloured according to the species colour scheme on the bottom right. **e**, Gas chromatogram of leaves volatiles from *Z. armatum* and *Z. bungeanum*. **f**, Detection of resistance of *Z. armatum* and *Z. bungeanum* to chewing insects (*Helicoverpa armigera*). The green, red and blue arrow represents the leaves of *Z. armatum*, *Z. bungeanum* and other test species (words in white) respectively. Scale bars, 1 cm.

Figure 6 Genomic basis of adaptive evolution in Chinese pepper. **a**, Expansion of genes involved in stress resistance and flavonoid metabolism. **b**, Expression profiles of genes encoding enzymes involved in flavonoid biosynthesis. The gradient color for each gene represents the gene expression levels in root, leaf and fruit tissues of A subgenome of *Z. armatum*, B and C subgenomes of *Z. bungeanum*. The colored dot on the left of the heatmap indicates whether the gene is affected by TE (with or without TE inserted into the UTR, exon or intron region of the gene). The color is consistent with the species color label in the heatmap. Gray indicates that the gene is not affected by TE. **c**, Liquid chromatogram of flavonoid metabolism from leaf of *Z. armatum* and *Z. bungeanum*. Principal component (PC)

analysis (d) and content (e) of flavone related metabolites in representative cultivars of *Z. armatum* and *Z. bungeanum* in different cultivation regions.

Figure 7 **The graphic illustration of evolution.** a, Square part represents the evolutionary characteristics of angiosperm in the mid-later Cretaceous and early Cenozoic, mainly including morphogenesis of floral organ, biosynthesis of kairomones/synomones, co-evolution with insect pollinators, and rapid growth under warmly and equable climates. b, Round part represents the evolutionary characteristics of Chinese pepper in the mid-later Cenozoic, mainly including the WGDs events (genomic evolution), resistance to abiotic and biotic stress (phenotypic innovation) and severe environmental factor. The taller plant with red fruit represents *Z. bungeanum*. Plant with green fruit represents *Z. armatum*. The shorter height suggests the narrow distribution, which shaped by cold stress.

Table 1 Summary of the genome assemblies for *Zanthoxylum armatum* and *Zanthoxylum bungeanum*

	Zanthoxylum armatum (2n = 4x = 132)			Zanthoxylum bungeanum (2n = 4x = 132)			Zanthoxylum armatum	Zanthoxylum bungeanum
	Monoploid	Subgenome A1	Subgenome A2	Monoploid	Subgenome B	Subgenome C	(Wang, M. et al)	(Feng, S. et al)
Sequencing platform	PacBio Sequel II			PacBio Sequel II			PacBio Sequel	PacBio Sequel
Genome sequencing depth (x)	260	-	-	260	-	-	114	100
Genome sequencing depth HiC	106	-	-	106	-	-	100	109
Estimated genome size (Gb)	4.4	-	-	4.6	-	-	3.1	4.43
Assembly								
Assembly Strategy	Canu+LACHESIS			Canu+LACHESIS			FALCON+LACHESIS	NextDenovo+ALLHiC
Number of scaffolds	148	33	33	173	33	33	14,619	332
Sequenced genome size (Gb)	4.39	2.21	2.06	4.63	1.96	2.48	2.64	4.23
Number of contigs > 100 Kb (%)	99.9%	100.0%	100.0%	96.5%	100.0%	100.0%	27.3%	-
Contig N90 (Mb)	1.01	1.02	1.03	3.06	2.83	3.93	0.05	-
Contig N50 (Mb)	3.31	3.34	3.23	10.74	9.77	13.67	0.34	0.41
Number of scaffolds > 100 Kb (%)	99.3%	100.0%	100.0%	96.5%	100.0%	100.0%	3.2%	-
Scaffold N90 (Mb)	47.43	48.63	47.33	43.23	47.57	44.55	0.16	-
Scaffold N50 (Mb)	68.34	70.14	65.07	65.42	58.90	92.33	71.48	74.18
Longest scaffold (Mb)	110.70	110.70	103.05	135.00	127.02	135.00	104.82	119.53
GC content (%)	36.41	36.39	36.45	36.09	36.36	35.89	36.29	36.81
Repetitive sequences (%)	82.40%	-	-	83.94%	-	-	80.13%	89.00%
Annotated protein-coding genes	65,195	32,942	28,618	68,202	31,074	28,596	55,355	74,307
BUSCO completeness of assembly (%)	98.1	90.0	84.7	98.5	96.5	92.4	94.6	97.59
BUSCO completeness of annotation (%)	98.2	87.0	81.7	98.7	93.7	90.8	91.7	-

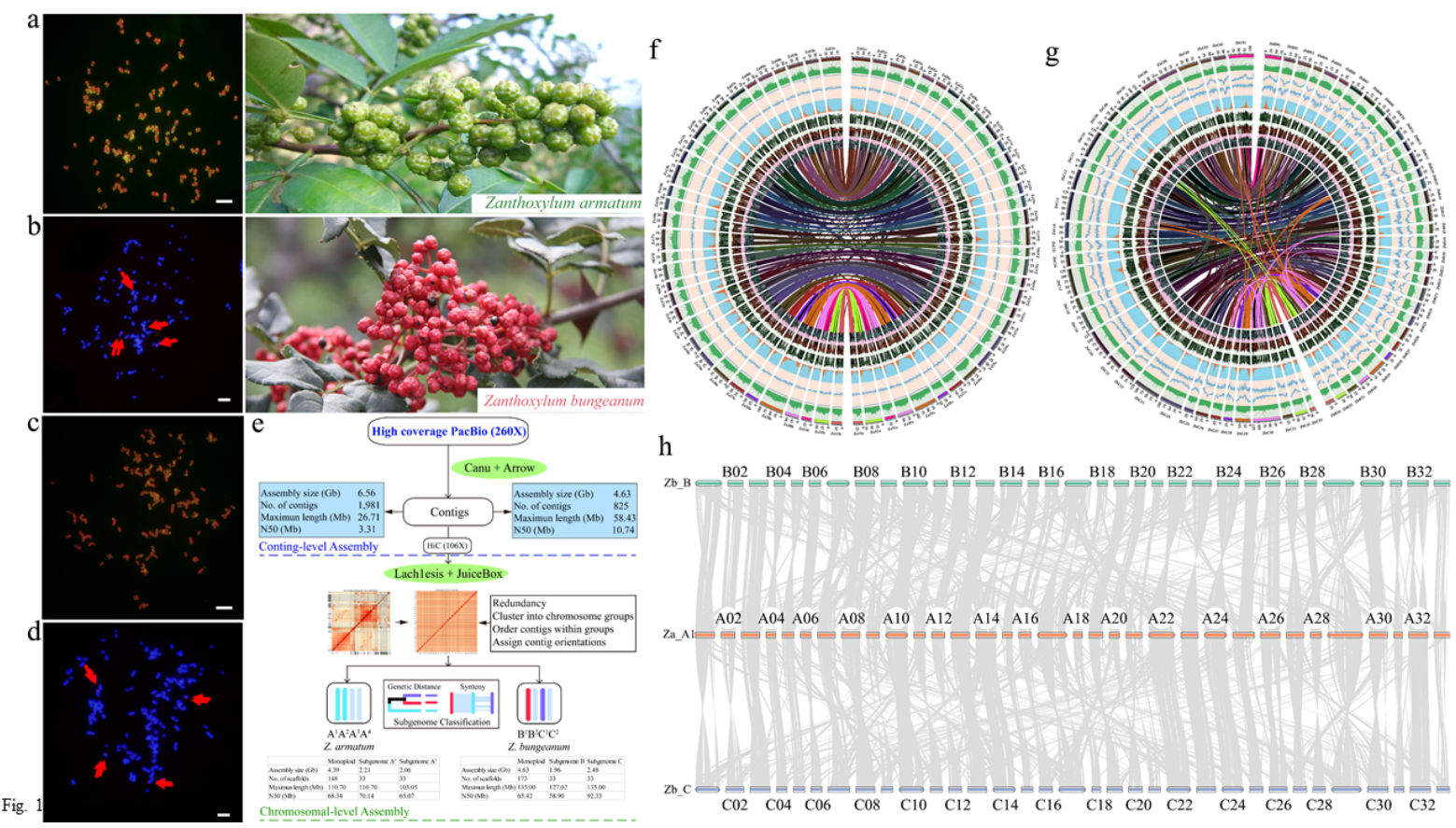


Fig. 1



Fig. 2

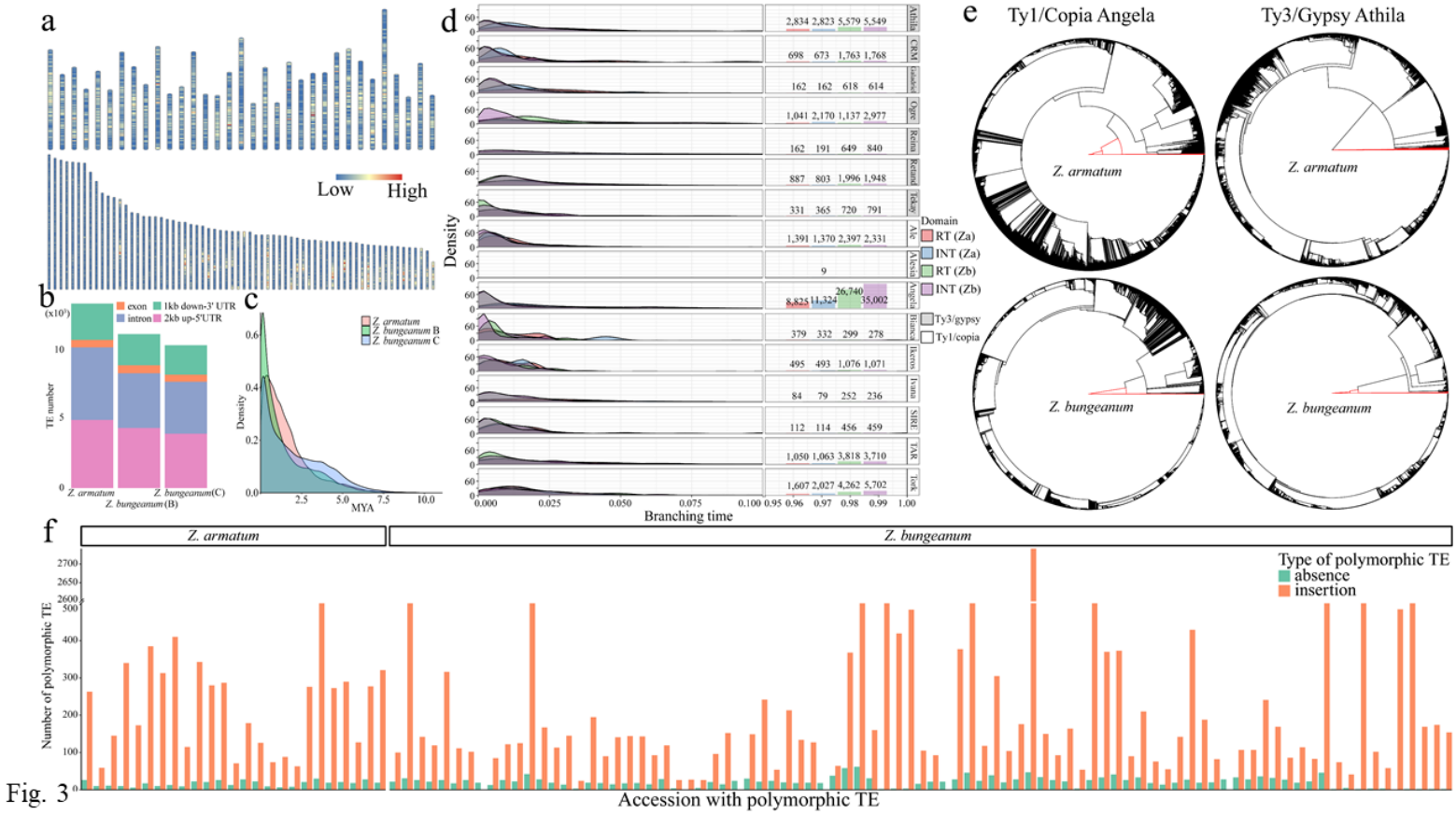


Fig. 3

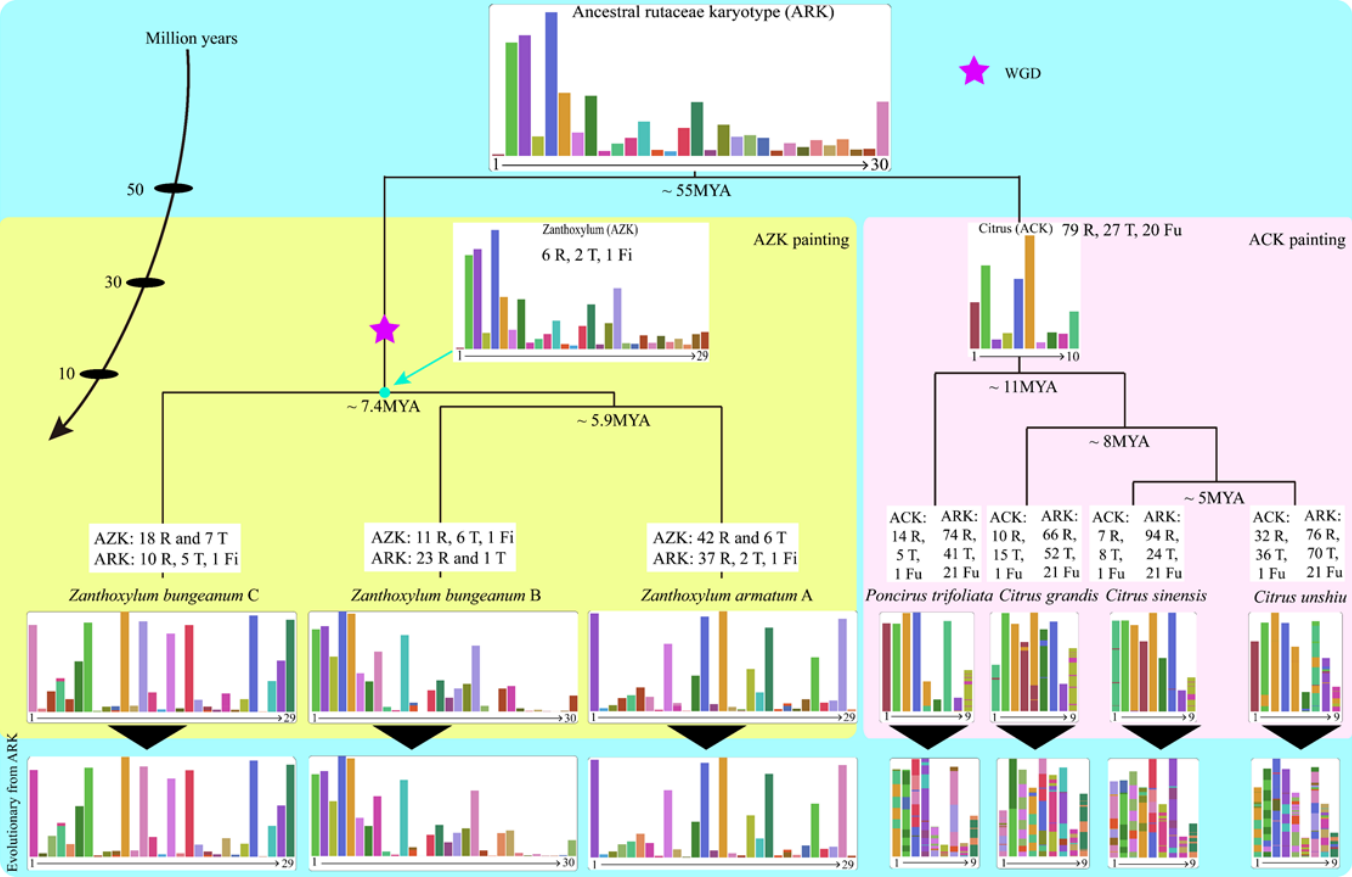
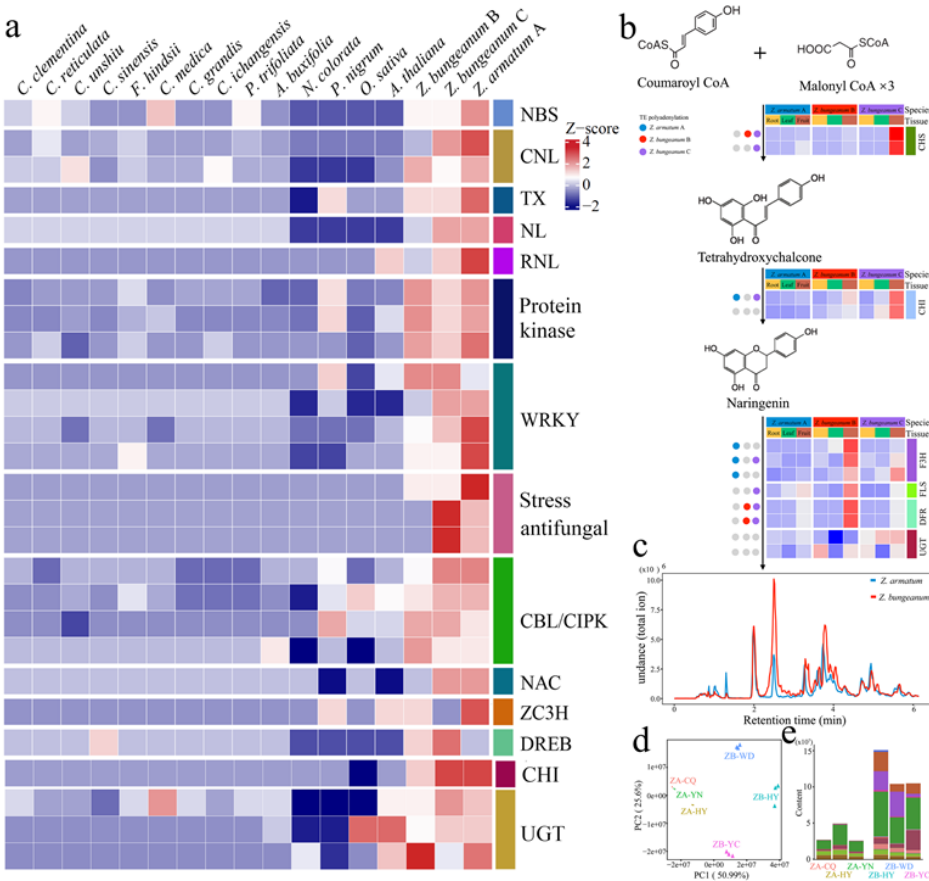


Fig. 4



Fig. 5

Fig. 6



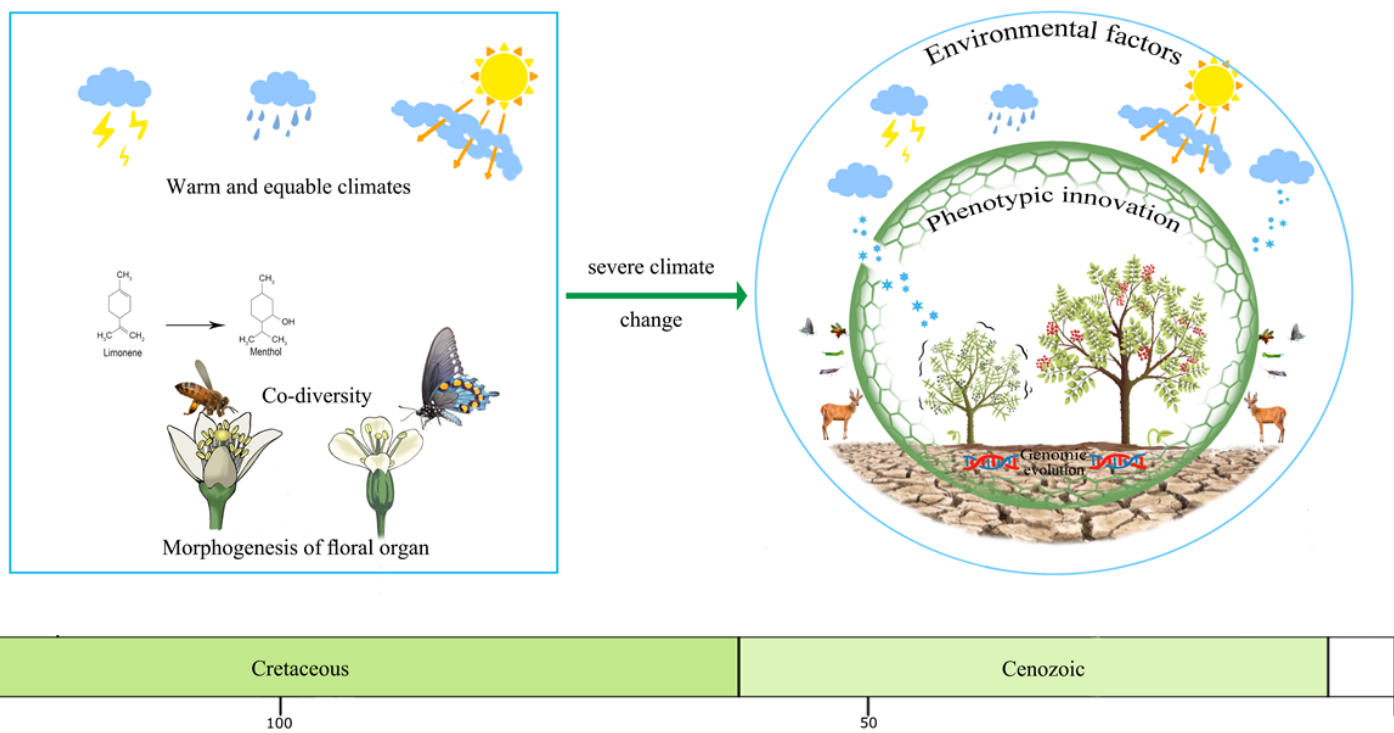


Fig. 7