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Genomic insights into longan evolution from a chromosome-level genome assembly and population analysis of longan accessions — Source link \square

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1	Research Article
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3	Genomic insights into longan evolution from a chromosome-level genome
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29 ABSTRACT

Longan (Dimocarpus longan) is a subtropical fruit best known for its nutritious fruit 30 and has been regarded as a precious tonic and traditional medicine since ancient times. 31 32 High-quality chromosome-scale genome assembly is valuable for functional genomic 33 study and genetic improvement of longan. Here, we report a chromosome-level 34 reference genome sequence for longan cultivar JDB with an assembled genome of 455.5 Mb in size anchored to fifteen chromosomes, representing a significant 35 improvement of contiguity (contig N50=12.1 Mb, scaffold N50= 29.5 Mb) over a 36 37 previous draft assembly. A total of 40,420 protein-coding genes were predicted in D. *longan* genome. Synteny analysis suggests longan shares the widespread gamma event 38 39 with core eudicots, but has no other whole genome duplications. Comparative genomics 40 showed that D. longan genome experienced significant expansions of gene families 41 related to phenylpropanoid biosynthesis and UDP-glucosyltransferase. Deep genome 42 sequencing analysis of 87 longan accessions identified longan biogeography as a major contributing factor for genetic diversity, and revealed a clear population admixture and 43 44 introgression among cultivars of different geographic origins, postulating a likely 45 migration trajectory of longan overall confirmed by existing historical records. The chromosome-level reference genome assembly, annotation and population genetic 46 resource for *D. longan* will facilitate the molecular studies and breeding of desirable 47 longan cultivars in the future. 48

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50 Keywords: *Dimocarpus longan*, reference genome assembly, phenylpropanoid, gene
51 flow, population genomics

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53 INTRODUCTION

Longan (Dimocarpus longan Lour.), also known as dragon's eveball and closely related 54 to lychee, is a tropical/subtropical evergreen fruit tree in Sapindaceae family with a 55 diploid genome¹ (2n = 2x = 30) It is an important economic fruit tree making great 56 contribution to the rural economic development in tropical and subtropical areas. It is 57 regarded as a precious tonic and traditionally used as a medicinal plant with rich 58 pharmaceutical effects from many parts of the plant, mainly fruits. The main functional 59 60 metabolites of longan include polysaccharides, polyphenols, flavonoids and alkaloids with anti-oxidative and anti-cancer activities². So far the biosynthetic pathways for 61 these metabolites in longan remain elusive due to limited genetic and genomic 62 63 resources and technical difficulty of genetic transformation.

Given its high nutritional and economic values, longan was cultivated in many countries 64 around the world, such as China, Australia, Thailand, Vietnam and other countries^{3, 4}. 65 China has the largest longan cultivation area and highest production⁵, including 66 Guangdong, Guangxi, Fujian, Hainan and other regions in China⁶. According to 67 historical records, longan is native to South China and has been cultivated for more than 68 2000 years in China with rich germplasm resources^{7,8} and lots of wild resources found 69 in Yunnan and Hainan province^{9, 10}, from which longan was introduced to other South 70 Asian countries such as Thailand^{11, 12}. A previous study based on the differences of 71 pollen exine patterns of fourteen longan varieties supports Yunnan as the primary center 72 of longan origin, and Guangdong, Guangxi and Hainan as the secondary centers¹³. 73 74 Thailand and Vietnam varieties have close genetic relationships indicated by ISSR (Inter-simple sequence repeat) analysis¹⁴. Although some molecular markers have 75

76 revealed genetic differences among germplasms, the classification of longan varieties based on these markers has differed among studies, due to different markers, number 77 of varieties and classification methods being adopted^{15, 16, 17}. Additionally, the 78 79 reproduction of longan can be achieved by both inbreeding and crossbreeding which 80 both bear seeds normally, therefore making longan varieties with ambiguous genetic 81 background. Therefore, a resolved population structure of longan varieties and understanding of its genetic diversity require a large-scale phylogenomic study of 82 83 longan varieties around China and Southeast Asia based on high quality genome 84 assembly and population resequencing data analysis. The knowledge of the longan 85 genetic background and its migration history is also required to improve longan 86 breeding.

87

88 Variety breeding has always been important for improving longan production, typically targeting two main traits, size and sweetness of the fruit^{18, 19}. The breeding and 89 extension of excellent varieties can enhance the stress resistance of fruit trees, improve 90 the fruit quality and expand the planting area. At present, it is challenging and time-91 92 consumming to improve longan by biotechnological breeding due to its long juvenile 93 period and difficulty of genetic transformation, sexual hybridization has been the main approach for longan breeding²⁰. Marker-assisted selection (MAS), based on the 94 identification of genes or genomic components related to desired new traits, is an 95 effective biotechnological tool to promote early selection of hybrid progenies at 96 seedling stage^{21, 22}. So far, our knowledge about genetic mapping of longan is limited. 97 98 Guo et al. (2011) constructed a low-quality male and female genetic map, consisting of 243 and 184 molecular markers separately²³. Single nucleotide polymorphism (SNP) 99

100 markers based on restriction site associated DNA sequencing (RAD-seq) was constructed for quantitative trait loci (QTL) identification by using hybrid progenies F₁ 101 and two parents as materials based on a draft genome sequence of *D. longan* "HHZ"¹⁹. 102 A chromosome-level reference genome sequence and knowledge of the longan genetic 103 104 background would significantly facilitate the investigation of genotype-phenotype 105 association of longan germplasms and thus expedite the longan breeding program. Although a draft genome sequence of *D. longan* "HHZ" cultivar was available²⁴, the 106 107 assembly is essentially fragmented composed of 51,392 contigs with a contig N50 of 108 26kb.

109

110 Here, we produced a chromosome-level genome assembly for *D. longan* JDB cultivar 111 combining Illumina paired-end (PE), PacBio single molecule real-time sequencing and high throughput chromatin capture sequencing (Hi-C). We annotated the genome using 112 113 ab inito prediction, homolog evidence and multi-tissue transcriptomic data. In addition, we conducted population genome deep sequencing from a collection of 87 longan 114 115 accessions, followed by an in-depth analysis of population structure using high-quality 116 genetic variants. The analysis revealed the population genetic diversity of longan and demonstrated the population admixture and introgression among cultivars from major 117 118 longan growing areas. The genome assembly, annotations and genetic variants will be valuable to functional genomic studies as well as molecular breeding of *D. longan* for 119 120 improving the yield, fruit quality and exploiting its medicinal properties.

121

122 **RESULTS AND DISCUSSIONS**

123 Genome assembly and annotation

D. longan "JDB" cultivar originated from Fujian is planted in Longan Germplasm 124 Repository of Guangdong Province (Figure 1A-1D), and the fresh young leaves were 125 collected for genomic DNA isolation and sequencing. To generate chromosome-level 126 genome assembly for *D. longan*, we produced 184.4 Gb PacBio single molecule 127 sequencing reads (415x coverage), 25.3Gb (56x coverage) Illumina paired-end (PE) 128 129 reads, and 57.6Gb (127x coverage) chromosome conformation capture (Hi-C) Illumina read pairs (Supplementary Table 1). We estimated the genome size of *D. longan* cultivar 130 131 JDB as 474.98 Mb with a heterozygosity rate of 0.36% via k-mer frequency analysis 132 using Illumina PE reads (Figure 1E). High-quality PacBio single-molecule sequencing reads were used to assemble the *D*. longan genome by using $Canu^{25}$, followed by 133 polishing contigs using Illumina PE reads by using $Pilon^{26}$, which yielded a draft 134 genome assembly of 455.5Mb (Table 1). Next, Hi-C paired-end reads were used to 135 anchor the PacBio assembled contigs to chromosomes with $Juicer^{27}$ and $3D-DNA^{28}$. 136 137 The final *D. longan* JDB genome assembly of 455.5Mb covers 95.90% of the estimated genome size (474.98Mb) and 98.7% of sequences were anchored onto 15 chromosomes 138 139 (Figure 1F) with contig and scaffold N50 of 12.1Mb and 29.6Mb, respectively (Table 140 1). Thus, this longan genome assembly represents a significant improvement over the highly fragmented D. longan HHZ genome assembly (contig N50 0.026 Mb) 141 previously released²⁴. Genome completeness was assessed using the plant dataset of the 142 Benchmarking Universal Single Copy Orthologs (BUSCO) database v1.22²⁹, with e-143 value < 1e-5. BUSCO evaluation revealed the completeness of 98.1% for our *D. longan* 144 genome assembly (88.4% single copy; duplicated copy 9.7%, 1.1% fragmented and 0.8% 145 missing) (Table 1, Supplementary Table 2). 146

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149 Figure 1: Chromosome-level genomic assembly of longan (Dimocarpus longan Lour.). (A-150 **D**): Photos of flower (**B**), fruit cluster (**C**), and fruit section (**D**) of longan cultivar JDB. (**E**) 151 Kmer frequency distribution analysis for JDB genome based on Illumina paired-end reads. (F) 152 Overview of D. longan genome. Track a to i: chromosomes, GC-content, density of Gypsy LTR 153 (long terminal retrotransposons), density of Copia LTR, density of protein-coding genes, SNP 154 density, Indel density, distribution of secondary metabolic gene cluster (predicted using 155 plantismash), syntenic blocks (color ribbons). The density statistics is calculated within 156 genomic windows of 150kb in size.

157

We next performed genome annotations by using the BRAKER2 pipeline combining 158 159 evidences from ab initio prediction, protein homologs and multi-tissue (root, shoot, leaf 160 and fruit) transcriptome sequencing data. The genome annotation pipeline predicted a total of 40,420 protein-coding genes and 2,555 non-coding RNAs for D. longan, 161 162 respectively (Table 1). Longan genome has an overall guanine-cytosine (GC) content of 34 % and gene density of 89 genes per Mb (Supplementary Table 2). About 89.0 % 163 genes have been annotated with NR (non-redundant protein sequence database) and 164 165 84.6 % genes with KEGG (Kyoto encyclopedia of genes and genomes) terms

 of which 54.9% and 25.4% are long terminal repeat retrotransposons (LTRs) and DNA transposons respectively. Two major LTR subtypes, LTR-<i>Copia</i> (179.64 Mb) and LTR- <i>Gypsy</i> (66.18 Mb) represent 8.55% and 15.53% of the longan genome, respectively (Supplementary Table 4). 	166	(Supplementary Table 3). Repetitive elements make up 41.7 % of <i>D. longan</i> genome,
 transposons respectively. Two major LTR subtypes, LTR-<i>Copia</i> (179.64 Mb) and LTR- <i>Gypsy</i> (66.18 Mb) represent 8.55 % and 15.53 % of the longan genome, respectively (Supplementary Table 4). 	167	of which 54.9% and 25.4 % are long terminal repeat retrotransposons (LTRs) and DNA
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170 (Supplementary Table 4).	169	Gypsy (66.18 Mb) represent 8.55 % and 15.53 % of the longan genome, respectively
	170	(Supplementary Table 4).

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	Statistics	D. longan JDB	D. longan
		(this study)	Honghezi ²⁴
	Total number of contigs	250	51,392
	Assembly size (Mb)	455.5	471.9
Contig	Contig N50 (Mb)	12.1	0.026
	Contig N90 (Mb)	1.8	0.006
	Largest Contig (Mb)	31.1	0.17
	Total number of scaffolds	90	17,367
	Assembly size (Mb)	455.5	495.3
Scaffold	Scaffold N50 (Mb)	29.6	0.57
	Scaffold N90 (Mb)	22.3	0.12
	Largest scaffold (Mb)	46.6	6.9
	Number of genes	40,420	31,007
	Repeat content (%)	41.7	52.9
Annotation	Number of ncRNA	2,555	NA
	BUSCO (%)	98.1%	94%
	GC content (%)	43.9	33.7

172 Table 1. Statistics for *Dimocarpus longan JDB* genome assembly and annotations.

173

174 Comparative genomics and synteny analysis revealed longan whole genome175 triplication

176 Next, we performed intraspecies synteny analysis of *D. longan* genome to investigate

177 its genome evolution history. Intraspecies syntenic gene pairs in D. longan were 178 identified using MCScanX, which supported the presence of a whole genome triplication (WGT) event in longan genome (Figure 2A). Distribution of synonymous 179 substitution rate (Ks) for the syntenic gene pairs also supported that the D. longan 180 181 genome experienced the WGT (Figure 2B). The 1:1 ratio of syntenic blocks between longan and grape (Vitis vinifera) indicated that the longan WGT was the same event as 182 the grape WGT (γ) event, and no other whole genome duplication occurred following 183 184 longan-grape divergence (Figure 2C). Furthermore, the 1:2 ratio of syntenic blocks 185 between longan and poplar (Populus trichocarpa) confirmed that a species-specific WGD occurred in poplar but did not happen in longan (Figure 2C). 186

187 To reveal the genome evolution and divergence of longan, we performed phylogenomic analysis of longan and thirteen representative angiosperm species including eight 188 189 Rosids (Citrus sinensis, Carica papaya, Arabidopsis thaliana, Theobroma cacao, P. 190 trichocarpa, Ricinus communis, Glycine max, V. vinifera), two Asteroids (Solanum tuberosum, Nicotiana attenuata), one monocotyledon (Oryza sativa) and a basal 191 192 angiosperm (Amborella trichopoda). Orthogroup (gene family) identification revealed 193 that these plants shared 7530 orthogroups, 137 of which are single-copy ones (Figure 194 3A; Supplementary Table 5). Particularly, we identified 1366 orthogroups unique to D. longan comparing to A. thaliana, C. cinensis, S. tuberosum and P. trichocarpa (Figure 195 3B). The multiple sequence alignment of 137 single-copy orthologs in 14 species were 196 197 concatenated and used for phylogeny construction followed by a divergence time estimation using MCMCTREE calibrated with fossil record time (Figure 3C). We found 198 that among the thirteen species, longan was phylogenetically closest to C. sinensis, 199 200 which, both belonging to Sapindales. shared a last common ancestor at around 67

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201 million years ago (Mya) that diverged from asteroids (*N. attenuata, S. lycopersicum*) at



around 125 Mya (Figure 3C).

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Poplar

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205 Figure 2. Comparative genomics and synteny analysis of *Dimocarpus longan*. (A) Whole 206 genome dot plot of *D. longan* showing intraspecies genome synteny based on syntenic gene 207 pairs. The pair of black circles connected by a straight line highlight the syntenic blocks 208 detected in *D. longan* genome, which corresponds to the whole genome triplication (γ event). 209 (B) Distribution of Ks (synonymous substitution rate) density for syntenic paralogs or orthologs 210 detected in pairwise comparisons among various plant genomes. (C) Karyotype macrosynteny 211 plots displaying the collinear relationships for different chromosomes among grape (Vitis 212 vinifera), longan (Dimocarpus longan) and poplar (Populus trichocarpa). The colored lines 213 highlight the syntenic blocks conserved among three species.

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215 Phylogenomics reveals gene family expansion for phenylpropanoid biosynthesis

216 enzymes and UDP-glucosyltransferases

217 Gene family contraction/expansion are the evolutionary forces that drive the rapid speciation and result in the diversification of plants³⁰. Gene family analysis suggested 218 longan genome has experienced 1474 expanded gene families and 2424 contracted gene 219 220 families (Figure 3A). KEGG (Kyoto Encyclopedia of Gene and Genomes) enrichment of expanded and contracted gene families (P < 0.05) showed that the 312 expanded 221 222 gene families were significantly enriched with "phenylpropanoid biosynthesis", "phenylalanine metabolism", "anthocyanin", "sesquiterpenoid and triterpenoid 223 biosynthesis", "monoterpenoid biosynthesis" (Figure 3D). Longan is rich in flavonoids 224 225 and polyphenols, with anti-cancer, anti-oxidant properties in leaf, flower, fruits, and seeds^{24, 31, 32, 33}, which are derived primarily through phenylpropanoid pathways. The 226 227 branches of phenylpropanoid metabolism produce end products such as flavonoids, hydrox-ycinnamic acid esters, hydroxycinnamic acid amides (HCAAs), and the 228 precursors of lignin, lignans, and tannins³⁴. The phenylpropanoid pathway is one of the 229 most extensively investigated specialized metabolic routes³⁵. The 97 expanded longan 230 231 phenylpropanoid biosynthesis genes were classified into seven gene families: phenylalanine ammonia-lyase (PAL, 5 members), peroxidase (POD, 38 members), O-232 methyltransferase (OMT, 3 members), glycosyl hydrolase family 1 (GH1, 26 members), 233 aldehyde dehydrogenase family (ADH, 18 members) and AMP-binding enzyme (4 234 members), beta-galactosidase (BGL, 3 members) (Supplementary Table 6). They 235 participated in the biosynthesis of P-hydroxy-phenyl lignin, quaiacyl lignin, 5-hydroxyl-236 guaiacyl lignin and syringyl lignin, which are precursors of longan. It was speculated 237 238 that lignins were involved in the longan speciation as a major component of certain

plant cell walls³⁶. The presence of structural lignins can provide physical barriers 239 preventing the pathogen from entering the plant tissues³⁷, and required for mechanical 240 support for plant growth and facilitate the long-distance transportation of water and 241 nutrients³⁸. In these protection processes, key enzymes of phenylpropanoid and lignin 242 pathway were PAL, POD and PPO³⁹. PALs, the first enzyme in the phenylpropanoid 243 biosynthetic pathway, the majority in longan genome were expressed at the higher 244 levels in the roots, leaves and stems, none PAL was highly expressed in the green fruits 245 246 (Supplemental Figure 1) consistent with previous report²⁴. Among the 38 PODs in longan genome, 28 showed differential expression in four major tissues (leaves, stems, 247 roots and fruits) (Supplemental Figure 2). A previous longan genome study revealed 248 non-expanded structural genes involved in phenylpropanoid, and flavonoid pathwavs²⁴. 249 250 which dismatched with our result as expanded phenylpropanoid biosynthesis pathway. However, only PAL in phenylpropanoid pathway was studied, the other six gene 251 252 families as mentioned above were not studied in the past because of their nontissue-253 specific expression.

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Figure 3. Phylogenomic genomics of *Dimocarpus longan*. (A). Summary of gene family
clustering of *D. longan* and 13 related species. Single copy orthologs: 1-copy genes in ortholog
group. Multiple copy orthologs: multiple genes in ortholog group. Unique orthologs: speciesspecific genes. Other orthologs: the rest of the clustered genes. Unclustered genes: number of
genes out of cluster. (B). Comparison of orthogroups (gene families) among six angiosperm
species including *D. longan* (longan), *A. thaliana* (Arabidopsis), *C. sinensis* (citrus), *S.*

263 tuberosum (potato), P. trichocarpa (poplar) and O. sativa (rice). (C). Phylogenetic relationship 264 and divergence time estimation. The number of gene family expansion and contraction was 265 indicated by red and blue number, respectively. (D). Bubble plot summarizing the most 266 significantly enriched KEGG (Kyoto Encyclopedia of Genes and Genomes) terms associated 267 with D. longan expanded gene families. X-axis is the log10 transformed p-value. The size of 268 bubble is scaled to the number of genes. The color scale represents the scale of odds ratio in 269 observed versus expected (genomic background) number of genes annotated with specific 270 KEGG terms. (E). A phylogenetic tree of UGTs (UDP-glucosyltransferase) in three 271 angiosperms including D. longan.

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InterPro (IPR) protein domain enrichment analysis showed that the expanded gene 273 274 families are significantly enriched with IPR domains such as UDP-glucosyltransferase (UGT) and Cytochrome P450s (Supplemental Figure 3). To cope with biotic and abiotic 275 stresses and interact with ecological factors for development, plants have evolved 276 exquisite mechanisms for the biosynthesis of secondary metabolites, through acylation, 277 methylation, glycosylation, and hydroxylation^{40, 41}. UGTs are key enzymes for 278 glycosylation, which can stabilize and enhance the solubility of small molecular 279 metabolites in order to maintain intracellular homeostasis^{42, 43}. Most of the compounds 280 synthesized by the phenylpropanoid pathway can be glycosylated by 281 glycosyltransferases⁴⁴. For example, UGTs were involved in the glycosylation of 282 volatile benzenoids/phenylpropanoids⁴⁵, and also monoterpene linalool⁴⁶, a strawberry 283 aroma 4-hydroxy-2,5-di-methyl-3(2H)-furanone⁴⁷ etc. A total of 215 UGTs were 284 285 identified in longan genome (Supplementary Table 7), more than in Arabidopsis (107), C. grandis (145), V. vinifera (181), but fewer than in apple $(241)^{48, 49, 50}$. UGTs 286 participate in multiple plant development and growth processes, including plant defense 287 responses^{51, 52}. It has been known for a long time that phenylpropanoid metabolism 288

plays important roles in resistance to pathogen attack^{53, 54}. A new mechanism of 289 phenylpropanoid metabolites reprogramming affecting plant immune response through 290 UGT has been revealed⁵⁵. In order to explore the evolutionary relationships of plant 291 UGT families, the phylogenetic tree was constructed based on the longan and other 292 plant UGT protein sequences, including Arabidopsis, Citrus (Figure 3e). All 115 293 294 expanded UGT members were divided into 10 phylogenetic groups. During the evolution of higher plants, the five phylogenetic groups A, D, E, G, and L appeared to 295 296 expand more than others, although the number of genes found in these groups varies widely among species⁵⁰. In longan, five phylogenetic groups A, D, H, I, and L expanded 297 more than the other groups, whereas no expanded longan UGTs were found in group G. 298 299 The number of longan UGTs in group D (31 UGTs, UGT73) and group I (19 UGTs, 300 UGT83) was significantly increased compared to those in Arabidopsis and Citrus. A group D member UGT73C7 was reported to mediate the redirection of phenylpropanoid 301 302 metabolism to hydroxycinnamic acids (HCAs) and coumarin biosynthesis under biotic stress, resulting in SNC1-dependent Arabidopsis immunity⁵⁵. In group I, number of 303 304 UGTs was highest compared to other fruits such as peach (5 UGTs), apple (11 UGTs) 305 and grapevine (14 UGTs). UGT83A1 (GSA1) was required for metabolite reprogramming under abiotic stress through the redirection of metabolic flux from 306 lignin biosynthesis to flavonoid biosynthesis and the accumulation of flavonoid 307 glycosides, which coordinately confer high crop productivity and enhanced abiotic 308 stress tolerance⁵⁶. In addition, the number of longan UGTs in groups H (14 UGTs, 309 310 UGT76) was reduced relative to Arabidopsis (21 UGTs). Transcript abundances of UGTs in different tissues were analyzed using RNA-seq data. Among longan UGTs, 96 311 312 UGTs were differentially expressed in longan. Additionally, four (accounting for 4.2%),

fourteen (14.6%), and ten (10.4%) UGTs were uniquely expressed in leaf, root, and fruit respectively (Supplemental Figure 4, Supplementary Table 8). The functions of the significantly expanded gene families highlighted the potential roles of these secondary metabolic enzymes to the longan genome evolution and adaptations.

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318 Tissue-specific expression of terpene biosynthesis genes in roots

Gene clustering is often associated with biosynthetic pathways for many plant natural 319 320 products. Therefore, we sought to identify gene clusters in the assembled longan 321 genome that may encode potential secondary metabolic pathways. Genome mining using *Plantismash* pipeline identified 29 secondary metabolic gene clusters in longan, 322 323 including 21 putatively involved in biosynthesis of alkaloids, saccharide and terpenes 324 (Supplementary Table 9). Tissue-specific transcriptome analysis showed that these gene clusters are expressed in various longan tissues (Supplementary Table 10). In the longan 325 326 genome, Chr3 contains four gene clusters associated with putative terpene biosynthesis (Supplementary Table 9). It has been reported that CYP450s play critical roles in 327 terpenoid skeleton modification and structural diversity^{57, 58}. CYP450 enzymes 328 329 involved in the terpenoid biosynthesis of pharmaceutical plants were mainly classified in three clans⁵⁹: CYP71, CYP85, CYP72. We found a longan gene cluster with 13 330 CYP450s on Chr3, which showed root-specific co-expression of eight CYP450 genes 331 (Supplementary Figure 5) within the Clan CYP71 (Supplementary Table 7). CYP450 332 superfamily was the second expanded gene family by IPR enrichment analysis. The 333 longan CYP450s (435) accounted for 1.1% of longan genes (Supplementary Table 7), 334 much higher than in Arabidopsis⁶⁰ (244), and grape⁶¹ (236). 335

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16

Root or phloem of longan has been used to treat filariasis, leucorrhea and other diseases 337 as traditional Chinese medicine. In the past, lots of metabolomics research focused on 338 longan leaf and fruit, whereas metabolic profiles of longan root were not clear. Terpenes 339 were chemical compounds responsible for plant's special odor and flavor profile⁶². 340 Puspita et al. (2019) reported that longan leaf ethanol extracts contained flavonoids and 341 triterpenoids⁶³. However, many of non-volatile terpenes were exuded from plant roots⁶⁴, 342 where they serve as the first line of plant defense and mediate below-ground 343 344 interactions between plants and other organisms. Therefore, the root-specific expression of a putative terpene biosynthesis gene cluster makes the terpene 345 accumulation and exudation much more effectively in longan roots, playing a role in its 346 347 biodefense against soil pathogens or herbivores.



348

349 Figure 4. Population structure and admixture analysis of Dimocarpus longan. (A). 350 Sampling localities of seven populations of *D. longan*, where red triangles distinguish the 351 sampling locations. (B). A neighbor-joining phylogenetic tree of all individuals of *D. longan* 352 was constructed using SNPs. The artificial breeding individual was marked with red dots inside. 353 Colors represent different geographic groups. (C). A biogeographical ancestry (admixture) 354 analysis of *D. longan* accessions with four ancestral clusters colored differently in the heatmap, 355 where each column represents a longan sample. (D). Distribution of *Fst* values (a measure of 356 genetic differentiation) between longan population from Thailand (Thai), Fujian (FJ) and

357 Guangdong (GD). (E). maximum-likelihood tree and migration events among seven groups of

358 *D. longan.* The migration events are colored according to their weight.

359

360 Population structure, migration and genetic admixture of longan cultivars

In the past, longan has been introduced among different populations frequently⁷. 361 Furthermore, longan can bear fruits by both inbreeding and crossbreeding²⁰. Lack of 362 363 reproductive barriers between native cultivars result in ambiguous genetic background of longan germplasms until now. To understand the longan genomic dynamics across 364 its current distribution range in southern China and southeast Asian countries, we 365 performed genome resequencing analysis of 87 accessions (Supplementary Table 11) 366 367 from five southern provinces in China: Guangdong, Fujian, Guangxi, Sichuan, Hainan, and three other countries Thailand, Vietnam and Australia, with an average sequencing 368 depth of 50×. Read mapping to longan reference genome and variant detection yielded 369 1,210,426 single nucleotide polymorphisms (SNPs), 204,991 insertions (INS) and 370 371 191,681 deletions. After filtering, 7,074,864 SNP loci were polymorphic (allele frequency > 0.05), among which 2,792,700 high-quality SNPs were used for 372 subsequent population genetic analyses. 373

374

Although Guangdong borders on Fujian (Figure 4A), the climate of the two provinces was largely different during longan growing season. After generations of planting and screening, different cultivation areas have formed their own longan variety characteristics and types. Using the genetic variant data, we analyzed the population structure within these longan cultivars using phylogenomic analysis and principal component analysis. Phylogenomic analysis clustered 87 longan samples into relatively distinct domestic Guangdong and Fujian groups after removal of artificial breeding

382 populations (Figure 4B). Three Sichuan cultivars were next to Fujian group and distant to GuangDong group. Notebly, two GuangDong cultivars, FLD and CPZ, were 383 clustered with Fujian group, probably because they come from eastern GuangDong 384 adjacent to Fujian. Guangdong cultivars are divided into two subgroups as "Shixai" 385 386 (SX) and "Chuliang" (CL) from central and western Guangdong, respectively (Figure 4B), also the two main cultivars widely grown in Guangdong and Guangxi. Consistent 387 with the phylogenetic tree, the principal component analysis of the 87 accessions 388 389 showed that Guangdong and Fujian cultivars were overall grouped separately, while 390 Thailand and Vietnam populations were distant to Chinese populations, when removing artificial breeding cultivars (Supplementary Figure 6). 391

392

393 To investigate the genetic background of longan from various regions, we performed 394 biogeographical ancestry (admixture) analysis based on high-quality SNPs and tested 395 it with ancestral group value (k) ranging from 1 to 10. With a choice of four ancestral groups (k=4) giving the smallest cross-validation errors (Supplementary Figure 7), the 396 397 admixture analysis discovered a distinct genetic structure within longan accessions of 398 different geographical origins. Longan cultivars from Fujian are composed of primarily two ancestral groups, whereas Guangdong, Guangxi and Sichuan cultivars contain 399 fractions of all four ancestral groups, indicating their more complex ancestry 400 backgrounds than Fujian ones (Figure 4C). The more similar ancestry composition 401 402 between eastern Guangdong and Fujian cultivars is accordant to the geographical closeness of the two growing regions, suggesting their common ancestral origin or a 403 possible exchange of cultivars between the two regions. By contrast, Thailand and 404 Vietnam cultivars overall have a simple composition with predominantly one ancestral 405

group, most likely shared with western Guangdong and Guangxi cultivars (Figure 4C).
Thailand cultivars were genetically more related to western Guangdong cultivars
(Figure 4B), but distant from Fujian cultivars. Consistent with this, we have also
detected a stronger genetic differentiation (measured in *Fst* value) between Thailand
and Fujian than between Thailand and Guangdong (Figure 4D). Notably, the Australian
cultivar has a genetic background resembling middle Guangdong cultivars, suggesting
it is likely a cultivar of middle-Guangdong origin introduced into Australia lately.

413

414 With the diverse ancestry backgrounds in these longan cultivars, we are curious about the migration history of longan germplasms and therefore investigated potential gene 415 416 flows among different growing areas due to such migration using Treemix analysis. 417 Given its reported origin in China, lots of wild longan resources are present in Yunnan and Hainan province of China^{9, 10}. Therefore, the Hainan cultivar was used as an 418 419 outgroup in this analysis. The Treemix analysis detected a migration event directed 420 from Hainan to Guangdong. There was the highest gene flow (migration weight 0.44) 421 between Sichuan and Fujian (Figure 4E). Gene flows were also detected from the Fujian, 422 Guangdong and Guangxi populations to Thailand with a high weight (migration weight 0.31) (Figure 4E). The detection of gene flows was consistent with longan migration 423 history on record. Longan was first cultivated in 'Ling-nan' district of China including 424 Guangdong, Guangxi and Hainan about 2000 years ago, recorded by painting of "San 425 Fu Huang". According to history records, longan was moved to northern China-Shaanxi 426 Province unsuccessfully, but was successfully introduced to Sichuan and then Fujian 427 with suitable climate conditions (Yang Fu, "Chronicles of the South", 1st century A.D.). 428 429 Taken together, our analysis results overall matched history records that there was gene

- 430 flow from Hainan wild germplasms to Guangdong, then a strong flow from Sichuan to
- 431 Fujian, and finally the gene flow from China to Thailand.
- 432

433 MATERIALS AND METHODS

434 Germplasm genetic resources

A 30-year-old *D. longan* tree cultivar named JDB from the Institute of Fruit Tree
Research at Guangdong Academy of Agricultural Sciences in China was used for
genome sequencing and *de novo* assembly in this study. Eighty-seven additional *D. longan* cultivars (Supplementary Table 11) that are widely grown in Southern China
and other countries were collected for genome resequencing.

440 **DNA and RNA isolation**

441 Longan cultivar JDB was planted in Longan Germplasm Repository of Guangdong Province. The fresh and healthy young leaves were collected, cleaned and used for 442 genomic DNA isolation and sequencing. Genomic DNA was extracted from young 443 444 fresh leaves of *D. longan* using the modified cetyltrimethylammonium bromide (CTAB) method⁶⁵. The concentration and purity of the extracted DNA were assessed using a 445 446 Nanodrop 2000 spectrophotometer (Thermo, MA, USA) and Qubit 3.0 (Thermo, CA, 447 USA), and the integrity of the DNA was measured using pulsed-field electrophoresis with 0.8% agarose gel. In addition, fresh leaves and other tissues (roots, shoots, young 448 449 fruits) of JDB cultivar were collected for RNA isolation and transcriptome sequencing. Total RNA was isolated with RNAprep Pure Plant Kit (Tiangen Biotech) according to 450 the manufacturer's instructions. The integrity and quantity of extracted RNA were 451 analyzed on an Agilent 2100 Bioanalyzer. For each tissue, three biological replicates 452 were prepared for sequencing. 453

454 Genome and transcriptome sequencing

DNA sequencing libraries were constructed and sequenced on the Illumina NovaSeq 455 6000 platform at 50x depth according to the manufacturer's protocols (Illumina). To 456 generate long-read sequencing reads for D. longan, DNA libraries for PacBio SMRT 457 458 sequencing were prepared following the PacBio standard protocols and sequenced on a Sequel platform. In brief, genomic DNA was randomly sheared to an average size of 459 20 kb, using a g-Tube (Covaris). The sheared gDNA was end-repaired using polishing 460 461 enzymes. After purification, a 20-kb insert SMRTbell library was constructed according 462 to the PacBio standard protocol with the BluePippin size-selection system (Sage Science) and sequences were generated on a PacBio Sequel (9 cells) and PacBio RS II 463 464 (1 cell) platform by Biomarker Technologies. Raw subreads was filtered based on read quality (≥ 0.8) and read length(≥ 1000 bp). For chromosome-level genome scaffolding, 465 Hi-C libraries were prepared from fresh leaves following protocol previously reported 466 ⁶⁶and sequenced on the Illumina HiSeq X Ten platform. DNA was digested with HindIII 467 enzyme, and the ligated DNA was sheared into size of 200-400bp. The resulting 468 469 libraries was sequenced by using Illumina NovaSeq 6000. For transcriptome 470 sequencing, RNA sequencing (RNA-seq) libraries were constructed using True-Seq kit (Illumina, CA), and sequenced using Illumina HiSeq X Ten platform. Illumina raw 471 472 reads were trimmed using Trimmomatic (v0.39) with parameters "LEADING: 10 TRAILING:10 SLIDINGWINDOW:3:20 MINLEN:36" to remove adapter sequences 473 474 and low quality reads, yielding a total of ~77.7 Gb clean RNA-seq data from four tissues. Genome assembly and evaluation 475

To estimate the genome size and heterozygosity level of *D. longan*, cleaned Illumina
PE reads were used for k-mer spectrum analysis using *kmergenie*⁶⁷ and *GenomeScope*

 $(v2.0)^{68}$ based on 21-mer statistics. PacBio SMRT reads were used for de novo genome 478 assembly by using *Canu* (V1.9)²⁵ pipeline with parameters "correctedErrorRate=0.045 479 corMhapSensitivity=normal 'batOptions=-dg 3 -db 3 -dr 1 -ca 500 -cp 50". Alternative 480 haplotig sequences was removed using *purge dups*⁶⁹ according default settings, and 481 only primary contigs were kept for downstream analysis. To correct the base-pair level 482 errors in raw assembly sequences, two rounds of polishing were conducted using high-483 gualitied Illumina DNA reads with *Pilon* $(v1.23)^{26}$. The longan contigs were further 484 anchored to chromosomes using *Juicer* $(v1.5.7)^{27}$ and *3D-DNA*²⁸ based on Hi-C contact 485 map, followed by manual correction using *Juicerbox* $(v1.11.08)^{70}$ to fix assembly errors. 486 The completeness of genome assembly was assessed by BUSCO $v1.22^{29}$ using 2121 487 488 eudicotyledons odb10 single copy genes. PacBio sequence reads and Illumina DNA reads were aligned to the genome sequences using *minimap2*⁷¹ and BWA⁷² respectively. 489

490 **Repetitive element annotation**

We used a combination of the *de novo* repeat library and homology-based strategies to 491 identify repeat structures. $TransposonPSI^{73}$ was used to identify transposable elements. 492 GenomeTools suite⁷⁴ (LTR harvest and LTR digest) was used to annotate LTR-RTs with 493 protein HMMs from the Pfam database. Then, a de novo repeat library of longan 494 genome was built using *RepeatModeler*⁷⁵, and each of the three repeat libraries was 495 classified with Repeat Classifier, followed by removing redundancy using 496 $USEARCH^{76}$ with \geq 90% identity threshold. Subsequently, the non-redundant repeat 497 library was analyzed using BLASTx to search the transposase database (evalue=1e-10) 498 and non-redundant plant protein databases (evalue=1e-10) to remove protein-coding 499 genes. Unknown repetitive sequences were further classified using CENSOR⁷⁷. Then, 500 501 the *de novo* repeat library was used to discover and mask the assembled genome with

502 $RepeatMasker^{78}$ with the "-xsmall -excln" parameter.

503 Prediction and annotation of protein-coding genes

For gene structure annotations, the RNA-seq data of four different tissues were aligned 504 to repeat-soft masked genome using $STAR^{79}$, which generates intron hints for gene 505 structure annotation. The structural annotation of protein-coding genes was performed 506 using *BRAKER2*⁸⁰, which integrates *GeneMark-ET*⁸¹ and *AUGUSTUS*⁸² by combining 507 the aligned results from *ab initio* predictions, homologous protein mapping, and RNA-508 509 seq mapping to produce the final gene prediction. The genes with protein length < 120amino acids and expression level < 0.5 TPM were removed. The tRNA genes were 510 identified by *tRNAscan-SE*⁸³ with eukarvote parameters. For rRNA, snRNA, miRNA 511 and other non-coding genes prediction, we used *INFERNAL*⁸⁴ software by search 512 against Rfam database⁸⁵. The contig-level genome sequences were used to blast against 513 plant plastid database from NCBI (https://ftp.ncbi.nlm.nih.gov/refseq/release/plastid/). 514 The organelle genome sequence identified was submitted to CHLOROBOX⁸⁶ website 515 to annotate and visualize. Predicted genes were assigned function by performing 516 BLAST against the NCBI non-redundant protein database with e-value threshold of 1e-517 10. In addition, a comprehensive annotation was also performed using InterProScan 518 (5.36-75.0)⁸⁷, which incorporates ProDom⁸⁸, PRINTS⁸⁹, Pfam⁹⁰, SMART⁹¹, 519 SUPERFAMILY⁹², PROSITE⁹³ database. Gene Ontology⁹⁴ identifiers for each gene 520 were obtained from the corresponding InterPro entry. $KAAS^{95}$ and $KOBAS^{96}$ were used 521 to search the KEGG GENES database for KO (KEGG Ontology) assignments and 522 generating a KEGG pathway membership⁹⁷. The stand-alone version of *plantiSMASH*⁹⁸ 523 was utilized to detect plant biosynthetic gene clusters in longan genome. 524

525 Comparative genomics analysis

526 Putative orthologship was constructed from two monocots, ten eudicots and Amborella trichopoda and longan proteome in this study. Only longest protein sequence was 527 selected as representative of each gene. Orthogroups were inferred by OrthoFinder 528 $(v2.4.1)^{99}$, as well as a STAG¹⁰⁰ species tree rooted using STRIDE¹⁰¹. The species tree 529 was used as a starting tree to estimate species divergence time using MCMCTREE in 530 paml (v4.9)¹⁰² package. Speciation event dates for Ananas comosus-Oryza sativa 531 (1.02~120 MYA), Populus trichocarpa-Ricinus communis (70~86 MYA), Arabidopsis 532 533 thaliana-Carica Papaya (63~82 MYA), and Glycine max-Citrus sinensis (98~117 MYA), which were obtained using *Timetree* database¹⁰³, were used to calibrate the 534 divergence time estimation. We conducted two independent MCMCTREE runs using 535 536 the following settings: burnin = 20000, sampfreq = 30, and nsample = 20000. The orthologous count table and phylogenetic tree topology inferred from the 537 *OrthoFinder* were taken into $CAF \acute{E}$ (v4.2)¹⁰⁴, which employed a random birth and death 538 539 model to estimate the size of each family at each ancestral node and obtain a familywise *p*-value to identify whether has a significant expansion or contraction occurred in 540 541 each gene family across species. Among expanded gene families, longan genes member 542 enriched in IPR002213 (UDP-glucuronosyl/UDP-glucosyltransferase) and IPR036396 (Cytochrome P450 superfamily) and their ortholog CDS sequences of A.thaliana and 543 C. sinensis genome were retrieved. Genes with protein length <300 amino acids were 544 removed. Multiple sequence alignment was conducted using MUSCLE (v3.8.1551)¹⁰⁵ 545 546 software. *IQtree* was used to constructed a maximum likelihood tree with parameters "-m MF". Tree file was loaded into the Interactive Tree of Life (iTOL) web server for 547 tree visualization and figure generation¹⁰⁶. 548

549 Transcriptomic analysis

After removing adapters and trimming low-quality bases, RNA-seq reads were mapped to the longan reference genome using $STAR^{79}$ with parameters "--alignIntronMax 6000 –align IntronMin 50" and then using *RSEM* tool¹⁰⁷ for transcripts quantification. Outliers among the individual experimental samples were verified based on the Person correlation coefficient, $r^2 \ge 0.85$. Differential expression analysis was performed using *DEseq2*¹⁰⁸ package. Genes were differentially expressed between two conditions if the adjusted p-values was <0.01 and fold change > 1.

557 Genetic variation detection

Genome resequencing data were mapped to chromosome-level genome assembly of 558 longan using *BWA-mem*⁷². *Bammarkduplicates* tool in *biobambam*¹⁰⁹ package was used 559 560 to mark and remove duplicate reads from individual sample alignments. Variant calling was performed using Freebayes¹¹⁰ with parameter "-C 5 --min-alternate-count 5 -g 561 10000" and then normalized with VT^{111} , filtered using vcffilter from vcflib¹¹² package 562 with parameters "QUAL / AO > 10 & SAF > 2 & SAR > 2 & RPL > 2 & RPR > 2 & 563 AF > 0.1". Only biallelic variants occurred in more than 90% of individuals were kept 564 565 and involved in further analysis.

566 **Population structure and history inference**

The vcf-format SNP set were transformed into binary ped-format using VCFtools¹¹³ and $PLINK^{114}$, and then *smartPCA*¹¹⁵ was used to conduct PCA analysis based on the data generated in the last step. High-quality SNP data were used to construct individual phylogenetic relationship with SNPhylo¹¹⁶ package. To estimate individual admixture assuming different numbers of clusters, the population structure and ancestry were investigated using *ADMIXTURE*¹¹⁷ based on all SNPs. An LD pruning step was performed with Plink¹¹⁷ with parameters "--indep-pairwise 50 10 0.1". We selected 574 ancestry clusters number ranging from 2 to 4. The population structure result was plotted with script downloaded from https://github.com/speciationgenomics. To study 575 the genetic relationship between longan population from different region, we computed 576 D-statistics. The calculation was performed with admixr¹¹⁸ package in the form of 577 (((Population1, Population2), Population3), HN), where Population1, Population2 and 578 Population3 represented longan from different lineage. Only combinations with 579 absolute Z-score value >3 were treated as confidential results. The demographic history 580 581 of longan was inferred using a hidden Markov model approach as implemented in pairwise sequentially Markovian coalescence¹¹⁹. We chose the default PSMC setting "-582 N25 -t15 -r5 -p 4+25*2+4+6" for all individual. To determine variance in Ne estimates, 583 584 we performed 100 bootstraps. We scaled results to real time estimates of generation time and mutation rate. We used synonymous substitution rate per synonymous site and 585 dated phylogeny tree as proxies for mutation rate estimation. 586

Syntenic blocks and reciprocal best hit orthologous pair were identified using 587 McScanX¹²⁰ with "--full --cscore=.99" parameters. Gene CDS were used as queries to 588 589 search against the genomes of the other plant genome sequences to find best matching 590 pairs. Given both CDS and protein sequence alignment of each gene pair, $PAL2NAL(v14)^{121}$ was subsequently used to perform codon alignment and 591 KaKs Calculator 2.0¹²² calculate Ka and Ks value under YN00 model. Gaussian 592 mixture models were fitted to the resulting frequency distribution of Ks values by means 593 of function density *Mclust* in the R *mclust* package (v5.3)¹²³. The Bayesian information 594 criterion was used to determine the best-fitting model for the data, including the optimal 595 number of Gaussian components as one. The formula r = D/2T, where D is the median 596 of Ks value, was used to estimate the neutral mutation rate. Mutation rate of 1.4×10^{-8} 597

- 598 per site per generation, and a constant generation time were assumed in this study to
- 599 convert coalescence generations into time-scale.

600 AUTHOR CONTRIBUTIONS

- 601 Project design and oversight: LG, JL and WQ; Sample collection and curation: DG and
- 602 SH; Conducting experiment and data analysis: JW, ZL and LG; Result interpretation:
- 603 LG, JL, JW, BL and WQ; Figure and table preparation: LG, JW and LZ; Manuscript
- 604 writing and revision: LG, JW, LZ, BL and WQ; Provide funding: JL and LG; All authors
- have read and proved the final version of this manuscript.

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614 CONFLICT OF INTEREST

615 The authors declare no conflict of interest.

616 SUPPLEMENTARY MATERIALS

617 Supplementary Figure 1: The heatmap of phenylalanine ammonia-lyase genes (PALs)

- 618 expressed in various longan tissues.
- 619 Supplementary Figure 2: The heatmap of peroxidase genes (PODs) expressed in
- 620 various *Dimocarpus longan* tissues.

- 621 Supplementary Figure 3: InterPro protein domain enrichment analysis of *Dimocarpus*
- 622 *longan* expanded gene families.
- 623 Supplementary Figure 4: The heatmap of UGTs genes expressed in various longan
- 624 tissues.
- 625 Supplementary Figure 5: The heatmap of CYP450 clustered-genes expressed in various
- 626 longan tissues.
- 627 Supplementary Figure 6: Principle component analysis of *Dimocarpus longan* samples
- 628 based on genotypes.
- 629 Supplementary Figure 7: Biogeographical ancestry analysis with group value K.
- 630 Supplementary Table 1: Sequencing statistics.
- 631 Supplementary Table 2: Summary of Illumina data for genome survey and genome632 polishing.
- 633 Supplementary Table 3: Gene function annotated by different databases.
- 634 Supplementary Table 4: Statistics of repetitive elements.
- 635 Supplementary Table 5: Comparison of genes in orthogroups between *Dimocarpus*
- 636 *longan* and 13 other species.
- 637 Supplementary Table 6: List of phenylpropanoid biosynthesis genes and their
- 638 expression level in different tissues.
- 639 Supplementary Table 7: List of different expressed IPR enriched gene families.
- 640 Supplementary Table 8: List of UGTs genes ID and their expression level.
- 641 Supplementary Table 9: The gene clusters found in *Dimocarpus longan* genome.
- 642 Supplementary Table 10: Tissue-specific transcriptome analysis of gene clusters
- 643 expressed in various longan tissues.
- 644 Supplementary Table 11: List of genome resequencing samples and their locations.

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