# 1 Comparative Population Genomics of Bread Wheat (Triticum

# 2 aestivum) Reveals Its Cultivation and Breeding History in China

- 3 Haofeng Chen<sup>1,2,7,8</sup>, Chengzhi Jiao<sup>3,8</sup>, Ying Wang<sup>1,8</sup>, Yuange Wang<sup>2,8</sup>, Caihuan Tian<sup>2</sup>,
- 4 Haopeng Yu<sup>1,2,4</sup>, Jing Wang<sup>2</sup>, Xiangfeng Wang<sup>5</sup>, Fei Lu<sup>1,6</sup>, Xiangdong Fu<sup>1,6</sup>,
- 5 Yongbiao Xue<sup>1,6</sup>, Wenkai Jiang<sup>3</sup>, Hongqing Ling<sup>1,6</sup>, Hongfeng Lu<sup>3</sup>\*, Yuling Jiao<sup>1,2</sup>\*
- <sup>6</sup> <sup>1</sup>College of Life Sciences, University of Chinese Academy of Sciences, Beijing 100049,
- 7 China.
- 8 <sup>2</sup>State Key Laboratory of Plant Genomics, Institute of Genetics and Developmental Biology,
- 9 Chinese Academy of Sciences, Beijing 100101, China.
- <sup>3</sup>Novogene Bioinformatics Institute, Beijing 100083, China.
- <sup>4</sup>West China Biomedical Big Data Center, West China Hospital/West China School of
- 12 Medicine, and Medical Big Data Center, Sichuan University, Chengdu 610041, China.
- 13 <sup>5</sup>Department of Crop Genomics and Bioinformatics, College of Agronomy and
- 14 Biotechnology, National Maize Improvement Center of China, China Agricultural University,
- 15 Beijing 100193, China.
- <sup>6</sup>State Key Laboratory of Plant Cell and Chromosome Engineering, Institute of Genetics and
- 17 Developmental Biology, Chinese Academy of Sciences, Beijing 100101, China.
- 18 <sup>7</sup>Present address: National Research Institute for Family Planning, Beijing, 100081, China
- 19 <sup>8</sup>These authors contributed equally to this work.
- 20 \*Correspondence to: <u>yljiao@genetics.ac.cn</u> (Y.J.); <u>luhongfeng@novogene.cn</u> (H.Lu)
- 21

# 22 Abstract

- 23 The evolution of bread wheat (*Triticum aestivum*) is distinctive in that
- 24 domestication, natural hybridization, and allopolyploid speciation have all had
- 25 significant effects on the diversification of its genome. Wheat was spread around
- 26 the world by humans and has been cultivated in China for ~4,600 years. Here,
- 27 we report a comprehensive assessment of the evolution of wheat based on the
- 28 genome-wide resequencing of 120 representative landraces and elite wheat
- 29 accessions from China and other representative regions. We found substantially
- 30 higher genetic diversity in the A and B subgenomes than in the D subgenome.

Notably, the A and B subgenomes of the modern Chinese elite cultivars were mainly derived from European landraces, while Chinese landraces had a greater contribution to their D subgenomes. The duplicated copies of homoeologous genes from the A, B, and D subgenomes were commonly found to be under different levels of selection. Our genome-wide assessment of the genetic changes associated with wheat breeding in China provides new strategies and practical targets for future breeding.

38

# 39 Main

40 Bread wheat (Triticum aestivum) differs from other major grain crops, such as maize 41 (Zea mays), rice (Oryza sativa), and barley (Hordeum vulgare), by having an 42 allohexaploid genome with six sets of chromosomes, two sets each from three closely 43 related ancestral species that formed the A, B, and D subgenomes. Archaeological and 44 genetic evidence indicated that wheat underwent two polyploidization events during 45 its domestication. The first occurred ~0.82 million years ago between two diploid 46 species, T. urartu (AA) and an unknown close relative of Aegilops speltoides (SS), 47 which produced the allotetraploid *T. turgidum* (AABB). The second polyploidization 48 occurred during cultivation around 8,000–10,000 years ago, when T. turgidum 49 crossed with another diploid grass, Ae. tauschii (DD), to form the ancestral bread wheat  $(T. aestivum, AABBDD)^{1-4}$ . In addition, a recent genome analysis suggested 50 51 that Ae. tauschii originated from a more ancient homoploid hybridization event between the A and B lineage ancestors<sup>5</sup>. The A, B, and D subgenomes comprise the 52 53 ~17-Gb allohexaploid bread wheat genome. Enormous genome sequencing efforts 54 have resulted in the recent publication of a reference genome sequence for wheat<sup>4,6-9</sup>, 55 which has greatly enhanced our understanding of the genome of this vital crop.

56 After its evolution in the Middle East and Mediterranean regions, bread wheat 57 gradually spread to the rest of the world and was domesticated for human use<sup>10</sup>. China

has been cultivating bread wheat for ~4,600 years<sup>11,12</sup>, and has been the largest wheatproducing country for more than two decades. Wheat has been under continuous artificial selection in the diverse ecological zones of China for thousands of years<sup>11,13</sup>; thus, the domestication and breeding of bread wheat in this country provides unique evolutionary insights into how its genome diversity was used and altered to meet the changing needs of the human population.

64 Recent advances in wheat genome sequencing have made pangenome studies possible<sup>14,15</sup>. To understand wheat genome diversity and evolution, we resequenced 65 66 120 landraces and elite bread wheat accessions at above a ten-fold coverage, and 67 employed the resequencing data to unravel the genealogical history of wheat 68 domestication. We identified the hybrid origins of various wheat cultivars and 69 landraces, especially those cultivated in China. Using these data, we began to 70 elucidate the genomic signatures that underlie human selection during the history of 71 wheat breeding.

72

#### 73 **Results**

#### 74 Characterization of molecular diversity

We selected 120 wheat accessions for our study, including 95 landraces and 25 75 76 cultivars. The 60 Chinese accessions are representative ones of a Chinese mini core 77 collection (Supplementary Fig. 1a and b), which covers the widest genome diversity of 23,090 Chinese landraces and cultivars<sup>16</sup>. The other accessions were selected based 78 79 on genotyping-by-sequencing results of 326 representative species collected 80 worldwide (Supplementary Fig. 1c). Notably, we selected 16 accessions from the 81 regions of the Fertile Crescent where wheat was first cultivated. Thus, the 82 resequenced accessions represent the widest genome diversity of both Chinese and 83 worldwide wheat populations. The geographic distributions of these accessions 84 included East Asia (China and Japan), West Asia, Central and South Asia, Europe, 85 and America (Fig. 1a and Supplementary Table 1).

86 A total of 21,676 Tb of sequence data was generated from the 120 accessions 87 using Illumina paired-end short-read technology, with an average read depth of 88 11.04× for each individual (Supplementary Table 2). The reads were mapped against 89 the IWGSC RefSeq v1.0 assembly of the wheat genome<sup>9</sup> to identify genomic variants, vielding a ~90% coverage of the genome by at least four reads for each wheat 90 91 accession. We detected a total of 70,172,600 single-nucleotide polymorphisms (SNPs) using the SAMtools software<sup>17</sup>, with an average density of 4.82 SNPs per kb. 92 93 Most of these SNPs were located in intergenic regions (94.76%), with only 1.43% 94 located in exonic regions (Table 1 and Supplementary Fig. 2). Of the SNPs located in 95 exonic regions, 36.62% were synonymous and 54.80% were non-synonymous, with a 96 non-synonymous/synonymous (N/S) ratio of 1.5. This N/S ratio is higher than was 97 previously reported for other plants such as sorghum (Sorghum bicolor; N/S ratio =  $(1.0)^{18}$ , rice (N/S ratio =  $1.2)^{19}$ , soybean (*Glycine max*; N/S ratio =  $1.47)^{20}$ , and 98 99 Arabidopsis thaliana  $(N/S \text{ ratio} = 0.83)^{21}$ . 100 We found that the A and B subgenomes harbor similar numbers of SNPs, with 101 the B subgenome (~32.95 M, 6.36 SNPs per kb) containing slightly more than the A 102 subgenome (~28.72 M, 5.82 SNPs per kb). By contrast, the D subgenome included 103 only ~7.91 M SNPs and had the lowest SNP density (2.00 SNPs per kb) (Table 1, Fig.

104 1c). A previous finding showed that low numbers of polymorphic loci in the D

105 subgenome may be a specific attribute of wheat, not of the D subgenome progenitor

106 *Ae. tauschii*<sup>22</sup>. In all three wheat subgenomes, the majority of SNPs were located in

107 intergenic regions, and non-synonymous SNPs were more common than synonymous

108 SNPs (the N/S ratios of A, B, and D subgenomes were 1.56, 1.59, and 1.28,

109 respectively; Table 1). In the wheat reference genome (IWGSC RefSeq Annotation

110 v1.0), the gene models were classified into high-confidence (HC) and low-confidence

111 (LC) genes based on their predicted sequence homology to gene products in the

112 public database. In the coding regions, 47.78% of the SNPs were located in HC genes.

113 These SNPs also had N/S ratios greater than 1 (Supplementary Table 3).

114 To measure the degree of polymorphism in the wheat genome, we measured 115 the nucleotide diversity parameters  $\pi$  and Watterson's  $\theta$ . The overall wheat genomic nucleotide diversity ( $\pi$ ) was  $1.05 \times 10^{-3}$ , and Watterson's  $\theta$  was  $0.84 \times 10^{-3}$ , which is 116 consistent with a previous report ( $\pi = 0.8 \times 10^{-3}$ )<sup>23</sup>. The nucleotide diversity parameter 117 for wheat was lower than for other major crops, including maize  $(\pi = 6.6 \times 10^{-3})^{24}$ , 118 rice  $(\pi = 2.29 \times 10^{-3})^{25}$ , and sorghum  $(\pi = 2.4 \times 10^{-3})^{26}$ . The  $\pi$  value of the cultivars 119  $(0.92 \times 10^{-3})$  was only slightly lower than that of the landraces  $(1.04 \times 10^{-3})$ . 120 Similarly, the Watterson's  $\theta$  value for the cultivars (0.84  $\times$  10<sup>-3</sup>) was slightly lower 121 than that of the landraces  $(0.87 \times 10^{-3})$ . The mean fixation index value ( $F_{ST}$ ) between 122 123 the cultivars and landraces was 0.03, suggesting limited population divergence 124 between the cultivars and landraces.

125 We then separated the data into the A, B, and D subgenomes to understand 126 their individual genetic diversities (Fig. 1b and Supplementary Table 4). In the 127 individual A, B, and D subgenomes, the nucleotide diversity parameters  $\pi$  and 128 Watterson's  $\theta$  were similar between the landraces and cultivars, indicating that the 129 modern wheat cultivars retained most of the genetic diversity of the landraces during 130 their domestication. The  $\pi$  and Watterson's  $\theta$  values each demonstrated that the A and 131 B subgenomes had similar nucleotide diversities, while the D subgenome had only 132 ~20% of the genetic diversity of the A or B subgenomes (Fig. 1c), consistent with a 133 previous report<sup>15</sup>. The heterozygous rates of all accessions were low (average: 134 0.1655%), reflecting the lack of cross-pollination, consistent with the cleistogamy of 135 wheat flowers. Among the three subgenomes, the D subgenome had a substantially 136 lower heterozygous rate than the A and B subgenomes (Supplementary Table 5), 137 consistent with its lower genetic diversity.

Tajima's *D* was calculated to assess whether the observed nucleotide
diversities showed evidence of deviation from neutrality. Some regions were
significantly different from zero, which indicates that they may be under sweep

141 selection (Fig. 1b). Of these, the D values from the A and B subgenomes were mostly 142 positive, whereas those of the D subgenome were generally negative (Fig. 1c and 143 Supplementary Table 4). Negative Tajima's D values indicate the presence of low-144 frequency SNPs in the D subgenome, while the positive Tajima's D values indicate a 145 predominance of intermediate-frequency SNPs in the A and B subgenomes. The 146 minor allele frequency (MAF) in the A, B, and D subgenomes significantly correlated 147 with the Tajima's D values (Supplementary Table 6). Because the three subgenomes 148 were expected to have experienced a similar evolution history in the allohexaploid 149 wheat after the second polyploidization event, the dramatic difference between D 150 subgenome and the A and B subgenomes indicates an asymmetric selection history 151 during domestication. Gene flow to bread wheat from wild and/or cultivated 152 tetraploid *T. turgidum* (AABB genome) is likely common, as suggested by the 153 identification of hybrid swarms between wild emmer wheat (T. turgidum subsp. 154 *dicoccoides*) and bread wheat $^{27,28}$ . By contrast, the barrier to gene flow from Ae. 155 tauschii (DD genome) to bread wheat is much more difficult to overcome. 156 Nevertheless, this difference may also contribute to the variations in the evolution 157 rates between the subgenomes.

#### 158 **Population structure**

159 To explore the phylogenetic relationships among the 120 wheat accessions, we 160 constructed a phylogenetic tree using the neighbor-joining algorithm, based on the 161 pairwise genetic distances between each accession determined using the SNP 162 information. The phylogenetic analysis clustered the wheat accessions into two 163 groups. Group 1 incorporated most of the European landraces, the West Asian 164 landraces (marked as "Origin"), a few South and Central Asian landraces, and the 165 majority of the East Asian cultivars (Fig. 2a). In group 2, most of the East Asian 166 landraces were clustered together alongside some of the West Asian landraces and the 167 majority of the South and Central Asian landraces (Fig. 2a). These two clearly defined 168 groups were further strengthened by the results of a principal component analysis

169 (PCA) and the Bayesian model-based clustering method (Fig. 2b). At K = 2, the group 170 1 accessions formed one cluster, while those of group 2 formed the other cluster. At 171 the optimal presumed number of ancestral populations (K = 3), the landraces near the 172 origin site (West, South, and Central Asian accessions) were separated from the main 173 groups. At K = 4, the West Asian and South and Central Asian accessions were 174 clearly divided. At K = 5, we obtained more refined clusters associated with the 175 geographic distributions (Fig. 2c and Supplementary Fig. 5). These findings were 176 consistent with the geographic documentation of wheat: from its domestication in the 177 Fertile Crescent, bread wheat was transported west to Europe and America, and east 178 to South and Central Asia and finally to East Asia via separate routes (Supplementary 179 Fig. 4)<sup>10</sup>. Our phylogenetic analysis, PCA, and clustering approaches revealed that 180 only a few Chinese cultivars are closely related to the Chinese landraces (Fig. 2a and 181 B). The population structure revealed that the Chinese cultivars are clearly related to 182 the European (and American) landraces when K = 2-5 (Fig. 2c). It is therefore evident 183 that the Chinese landraces contributed a minor portion of the genetic diversity of the Chinese cultivars. 184

185 To understand whether different diploid ancestors contributed similar amounts 186 of genetic diversity to the subgenomes, we further analyzed the population structure 187 of each subgenome separately. The A and B subgenome population structures 188 confirmed the evolutionary patterns observed for the whole genome (Supplementary 189 Figs. 6 and 7). In contrast, the D subgenomes of most Chinese cultivars were closely 190 related to the Chinese landraces in the phylogenetic tree analysis, which was further 191 supported by the results of the PCA and genetic structure analysis (Supplementary 192 Fig. 9). Taken together, the A and B subgenomes of the Chinese cultivars may mainly 193 come from European landraces and cultivars with a genetic admixture of Chinese 194 landraces, whereas the D subgenome contains more of a contribution from the 195 Chinese landraces with a lesser admixture of European landraces.

196	To investigate the differences in genetic diversity between the three
197	populations (Chinese cultivars, Chinese landraces, and European landraces), we
198	calculated the nucleotide diversities ( $\pi$ ) and found no significant difference among the
199	three populations. The diversity of Chinese cultivars was slightly higher than that of
200	the Chinese landraces, but slightly lower than that of the European landraces (Fig.
201	2d). We calculated the $F_{ST}$ values to investigate the population divergences, which
202	revealed that the divergence between the Chinese cultivars and European landraces
203	$(F_{ST} = 0.04)$ was smaller than the divergence of the Chinese cultivars and Chinese
204	landraces ( $F_{ST} = 0.07$ ). This analysis indicated a small population divergence (Fig.
205	2d).

206 We further analyzed the linkage disequilibrium (LD) of the three populations. The mean  $r^2$  between 0 and 1000 kb from the three populations was greater than 0.3, 207 208 suggesting that wheat has a longer LD decay than cultivated rice (123 kb for indica varieties) and cultivated maize  $(30 \text{ kb})^{29,30}$ . The mean  $r^2$  between 0 and 1000 kb of the 209 210 D subgenome decreased more rapidly than for the A and B subgenomes, suggesting a lower level of selection on the D subgenome (Fig. 2e). The extent of LD in the A, B, 211 212 and D subgenomes differed between the Chinese cultivars, Chinese landraces, and 213 European landraces. For the A subgenome, the LD decay of the Chinese landraces 214 was slightly higher than that of the European landraces, and significantly higher than 215 that of the Chinese cultivars. For the B subgenome, the Chinese landraces also had the 216 highest LD levels, with the Chinese cultivars and European landraces displaying 217 similar levels of LD decay. In the D subgenome however, the LD decay was similar 218 for all three populations. Different chromosomes had specific patterns of LD 219 (Supplementary Fig. 3), which may be correlated with differences in heterochromatin 220 levels<sup>31</sup> and selection pressure.

# Introgressions from Chinese landraces and European varieties in the Chinesecultivars

223 We used a TreeMix analysis to infer ancient gene flows. A maximum-likelihood 224 (ML) tree without migration events grouped the wheat accessions into seven clusters, 225 which were similar to the above population structure patterns. Furthermore, we 226 detected strong migration events in three clusters, namely the Chinese cultivars, 227 Chinese landraces, and European cultivars. There were strong gene flows from the 228 Chinese landraces to the Chinese cultivars, and from the European landraces to the 229 European cultivars (Supplementary Fig. 6a and b). We next individually analyzed the 230 three subgenomes. The ML trees for the A and B subgenomes of the accessions had 231 similar topological structures to the ML tree using the whole genomes (Fig. 3a). We 232 also found strong gene flows from the Chinese landraces to the Chinese cultivars and 233 from the ancient western landraces to the European landraces in both the A and B 234 subgenomes (Fig. 3a). When the D subgenome was considered, the accessions formed 235 a different ML tree structure, with Chinese landraces grouped with Chinese cultivars, 236 and clear gene flows from the European cultivars to the Chinese cultivars, and from 237 the South and Central Asian landraces to the European landraces (Fig. 3a).

238 To further elucidate the contributions from the European varieties and Chinese landraces to the Chinese cultivars, we used an identity score (IS) analysis<sup>32</sup> to scan 239 240 each chromosome. The IS analyses for the three groups (European varieties, Chinese 241 landraces, and Chinese cultivars) revealed that the Chinese landraces had more 242 similarity to the reference genome (Chinese Spring, a Chinese landrace) than the 243 European and Chinese cultivars, while the Chinese cultivars were more similar to the 244 European varieties than the Chinese landraces (Fig. 3b). Consistent with the results of 245 the population structure analysis, the IS heatmap showed that the European varieties 246 contributed more to the Chinese cultivars than did the Chinese landraces 247 (Supplementary Fig. 10). More differences were detected between the Chinese 248 landraces and the Chinese cultivars than between the European varieties and the 249 Chinese cultivars when using IS values less than 0.0025 as the threshold (Fig. 3c). 250 Specifically, 280,299 regions were detected, spanning 5.61 Gb (39.28% of the

genomes), in which the Chinese cultivars and European varieties were more similar
than the Chinese cultivars and Chinese landraces, while only 106,453 regions,
spanning 2.13 Gb (14.92% of the genomes) were identified in which Chinese
cultivars and Chinese landraces were more similar (Fig. 3d). Within most
chromosomes, we observed similar patterns of homology, although there were clear
variations (Supplementary Fig. 11, Supplementary Table 7a).

257 We were particularly interested in the many exceptionally large dispersed introgression regions identified in the genomes of the wheat population. Between the 258 259 Chinese landraces and the Chinese cultivars, the introgression regions are mainly 260 located in the A and B subgenomes, while those between the European varieties and 261 the Chinese cultivars were mainly located in the D subgenome. These introgression 262 regions usually show higher heterozygosity, low recombination rates, and long-range LD<sup>27</sup>; for example, an 8-Mb region in chromosome 5D of the Chinese cultivars is 263 264 enriched with SNPs from the European varieties (Fig. 3e). Within this region, the 265 Chinese cultivars are highly heterozygous and have low recombination rates, as well 266 as a significantly higher LD level (Fig. 3e). We focused on the mutation sites within 267 this candidate introgression region, which covers 346 genes (Supplementary Table 268 7b). Using a Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis, we 269 observed a significant enrichment of genes involved in pathways related to 270 photosynthesis within this region (Supplementary Table 7d), including genes involved 271 in photosystem I (TraesCS5D01G552400LC/PSAB, TraesCS5D01G552500LC/PSAB, 272 and TraesCS5D01G552600LC/PSAB) and photosystem II 273 (TraesCS5D01G550000LC/PSBA, TraesCS5D01G458100/PSBK, 274 TraesCS5D01G458200/PSBI, TraesCS5D01G550600LC/PSBA, 275 TraesCS5D01G550700LC/PSBC, TraesCS5D01G550800LC/PSBC, 276 TraesCS5D01G458300/PSBZ, and TraesCS5D01G458400/PSBM). Another 200-Mb 277 region on chromosome 6A of the Chinese cultivars is enriched with SNPs from the

278 Chinese landraces (Fig. 3f). This region also has higher heterozygosity and lower

recombination rates within these lines, and shows a significantly higher LD (Fig. 3f).
The 2,131 genes within this region (Supplementary Table 7c) were also enriched for
oxidative phosphorylation and ribosome pathway functions (Supplementary Table
7e). Taken together, our IS analysis revealed the detailed contributions of the
European landraces and Chinese landraces to the Chinese cultivars. Importantly, most
of these regions are genome-specific, suggesting the functional diversification of the
three genomes.

#### 286 Selective signals during diversification and modern breeding

287 To identify the genomic regions most affected by selection during wheat 288 diversification and modern breeding in China, we scanned the patterns of genetic 289 variation along the chromosomes on the basis of the 70.12 million SNPs. We 290 calculated the nucleotide diversity ratio ( $\theta_{\pi \text{ landrace}}/\theta_{\pi \text{ cultivar}}$ ) and genetic differentiation 291  $(F_{ST})$  between the cultivars and landraces. Long stretches of elevated  $F_{ST}$  were found 292 along chromosome 4A, which is known to contain structurally rearranged chromosomal regions<sup>33</sup>. After excluding these rearranged regions, the top 5% of 293 294 regions (~14.98 Mb) were considered putative sweep-selected sections, which 295 contained 842 genes (Supplementary Table 8a). A comparison among the 296 subgenomes indicated that selection in the D subgenome is substantially lower than in the A and B subgenomes, as indicated by the lower  $F_{ST}$  and nucleotide diversity ratio 297

298  $(\theta_{\pi \text{ landrace}}/\theta_{\pi \text{ cultivar}})$  for the D subgenome (Fig. 1b). The A subgenome had the most

selected regions (Supplementary Table 8b). We also used the XP-EHH<sup>34</sup> and XP-

300 CLR<sup>35</sup> methods to identify the regions under sweep selection, which respectively led

301 to the identification of 2.65 Mb (948 genes) and 16.63 Mb (2,168 genes) of the

302 putatively selected regions (Supplementary Table 8c and d). The combination of the

303 results from all approaches led to the identification of a total of 3,659 genes, the

304 majority of which were located on the A and B subgenomes (Supplementary Table

305 8f). The selected regions contain many previously reported quantitative trait loci

306 (QTL); for example, one on chromosome 6B and one on chromosome 7D both

associated with the length of the uppermost internode length<sup>36</sup>. Gene Ontology (GO)
and KEGG enrichment analyses revealed that the selected regions are enriched in
genes with decarboxylating activity and glycine catabolism, suggesting that the
carbon metabolic pathways were targeted during modern breeding. Additional
pathways, such as selenocompound metabolism, were also under selection
(Supplementary Tables 7g and h).

313 The allotetraploid nature of bread wheat raised the question of whether the 314 duplicate copies of the homoeologous genes are all under selection. Using the ordered 315 sets of homoeologous genes, we established the syntenic relationships of the selected 316 targets on the other subgenomes (Fig. 4b). This analysis indicated that some syntenic 317 regions shared the signature of selection; for example, the syntenic regions of 318 chromosomes 6A, 6B, and 6D were all under selection (Fig. 4b, Supplementary Table 319 8i). In the majority of cases however, we could not identify selection signatures in the 320 syntenic regions, suggesting that the homoeologous genes are generally under 321 differing selection pressures and are likely responsible for new functions.

322 To identify the high-confidence sweep-selected regions, we focused on the 323 overlapping regions commonly identified using the three above-mentioned 324 approaches, which contained 48 HC genes (Supplementary Table 8e). Chromosome 325 6B had the strongest selection signal. A sub-region of this selected region contains 326 TaNPF6.1-6B (Fig. 4c), an ortholog of the Arabidopsis nitrate transporter gene NRT1.1<sup>37,38</sup>. Within the same selected region, we also found TaNAC24, the ortholog 327 of a gene that confers heat and drought tolerance in rice<sup>39</sup>. In a neighboring sub-328 329 region, we found strong selection signals for TaRVE3 (which encodes a putative circadian clock and flowering time regulator)<sup>40</sup>, and *TaHDG2* (encoding a putative 330 stomata and epidermis patterning regulator)<sup>41</sup>, suggesting the existence of a selected 331 332 region regulating multiple traits on chromosome 6B (Fig. 4c). In another selected 333 region on chromosome 3A, we identified *TaHSP101*, which enhances tolerance to salt

and desiccation stresses<sup>42</sup>; *TaPTR7*, the ortholog of a rice NAC transcription factor

335 conferring salt tolerance<sup>43</sup>; and *TaPRO1*, a component on the actin cytoskeleton

- potentially related to grain size<sup>44</sup> (Fig. 4c). The majority of the selected features and
- 337 genes are currently poorly annotated and warrant further investigation.
- We analyzed the haplotype frequency of the annotated genes under sweep
- 339 selection, except *TaPTR7* as it contained no SNPs. In the landraces, the frequencies of
- the primary haplotypes of all selected genes were lower than in the cultivars (Fig. 4d
- and 4e). Different periods of wheat breeding history likely had diverse breeding goals.
- 342 To clarify the changes in the haplotypes of these genes over time, we screened the
- 343 SNPs of six genes in the selected regions (*TaAGL29*, *TaHDG2*, *TaKNAT6*,
- 344 *TaMADS21*, *TaRGI5*, and *WSD1-like*) for Chinese cultivars released between the
- 345 1940s and the 2000s (Supplementary Table 12b). For each gene, the frequency of the
- 346 favorable haplotype gradually increased over time (Fig. 4f).

## 347 Local adaption and related traits

348 Our phylogenetic analysis showed that landraces from geographically close regions 349 tended to be more genetically similar (Fig. 2). To understand the selection undergone 350 by the genes and corresponding traits of these genotypes during geographic 351 diversification and/or local breeding, we calculated the population differentiation 352 across different geographic groups.

353 The European landraces contributed the majority of the A and B subgenomes 354 of the Chinese cultivars (Supplementary Figs. 7 and 8). We compared these two 355 groups to identify the regions selected during the local adaptation of the Chinese 356 cultivars. Using the  $F_{\text{ST}}$ - $\theta_{\pi}$  approach, we identified 13.08 Mb of putatively selected 357 regions covering 729 genes (Fig. 5a). Over a dozen regions were similarly selected in 358 two or three of the subgenomes (Fig. 5b). We also applied the XP-EHH and XP-CLR 359 methods, which led to the identification of a 2.68-Mb putatively selected region 360 (covering 951 genes) and a 26.08-Mb putatively selected region (covering 3,232

361 genes), respectively (Supplementary Table 9). Finally, we excluded the selected 362 regions identified in our earlier comparison of all cultivars and landraces to focus on 363 local adaption. A total of 3,814 genes were identified using all three methods, with 51 364 genes being commonly detected (Fig. 5d and Supplementary Table 9e). Among the 365 regions commonly identified using all three approaches, we found that *TaPGIP1*, a polygalacturonase-inhibiting defense protein<sup>45</sup>; WRKY27, involved in disease 366 resistance<sup>46</sup>; *TaNAK*, a putative protein kinase involved in defense<sup>47</sup>; and *TaPHB2*, a 367 mitochondrial prohibitin complex protein<sup>48</sup>, were all located within a selected region 368 369 on chromosome 2A (Fig. 5c). The regions on chromosomes 2B and 2D syntenic to 370 this region were also under selection. Another selected region on chromosome 3A contains *TaPIN1*, required for auxin-dependent root branching<sup>49</sup>; *TaPrx*, related to 371 oxidative stress<sup>50</sup>; and *TaPEPR1*, involved in the innate immunity response<sup>51</sup> (Fig. 372 373 5c). A detailed haplotype frequency analysis indicated that the frequencies of the 374 primary haplotypes in the European landraces were lower than in the Chinese 375 cultivars for all six genes (Fig. 5e and f).

376 The West Asian landraces are the most closely related modern relatives of the 377 ancestral wheat population. We therefore compared the European and Chinese 378 landraces to the West Asian landraces to identify the regions selected during local 379 adaptation as wheat gradually spread to the west and the east. Using  $F_{ST}$ - $\theta_{\pi}$ , XP-EHH, 380 and XP-CLR analyses, we identified 22.75-Mb (covering 1,760 genes), 3.00-Mb 381 (covering 1,074 genes), and 21.04-Mb (covering 2,619 genes) putatively selected 382 regions between the European and West Asian landraces, respectively 383 (Supplementary Table 10). These selected regions overlap with a few reported QTLs, including two associated with flour quality<sup>52</sup>. We identified a significantly selected 384 385 region on chromosome 2B that contains TaNPF6.1-2B, another ortholog of the Arabidopsis nitrate transporter gene NRT1.1 (Supplementary Fig. 12a). The same 386 387 region also harbors *TaFBP7*, a gene related to temperature response<sup>53</sup>, and *TaANP1*, an oxidative stress responsive gene<sup>54</sup>. Whereas novel haplotypes for TaNPF6.1-2B388

emerged in the European landraces, the genotypes typically maintained one of the two *TaFBP7* and *TaANP1* haplotypes from the West Asian landraces (Supplementary Fig.
12b and c).

392 When comparing the Chinese landraces with the West Asian landraces, we 393 identified 8.94 Mb of putatively selected regions (covering 550 genes) using  $F_{\rm ST}$ - $\theta_{\pi}$ , 394 2.81 Mb (covering 1,027 genes) using XP-EHH, and 18.85 Mb (covering 2,392 395 genes) using XP-CLR (Supplementary Table 11). One of the genes identified in the 396 selected regions was TaOSH43 (Supplementary Fig. 13a), which regulates shoot meristem homeostasis and thus plant architecture and spike development<sup>55</sup>. A new 397 haplotype emerged and became dominant in the Chinese landraces (Supplementary 398 Fig. 13b and c). We also found that a flour-quality QTL<sup>52</sup> overlaps with a selected 399 400 region on chromosome 6B.

401

## 402 **Discussion**

403 Our analysis of 120 representative bread wheat accessions revealed a unique pattern 404 of genome changes that occurred during wheat adaption and modern breeding. 405 especially in China. The allopolyploidization of hexaploid bread wheat is expected to 406 have occurred under human cultivation, as bread wheat only exists in cultivated forms<sup>3,10</sup>. Consistent with this, bread wheat has a substantially lower genome diversity 407 408 than other crop and weed species (Fig. 1c and Table 1). Nevertheless, gene flow from 409 the allotetraploid T. turgidum (AABB) is likely to have contributed to the genetic diversity of the A and B subgenomes in bread wheat<sup>27,28</sup>, which means these two 410 411 subgenomes harbor more selective sweeps, as was demonstrated here. By contrast, the 412 barrier to gene flow from the diploid Ae. tauschii (DD) to bread wheat is much higher, 413 as suggested by the substantially lower nucleotide diversities in the D subgenome, 414 higher numbers of rare mutations, and fewer selective sweeps reported here.

415 Bread wheat has a long cultivation history and has been adapted to a wide 416 range of environmental conditions. The rich genetic diversity in cultivated accessions 417 and the lack of a wild bread wheat mean the gene flow among geographic regions is 418 potentially substantial. Indeed, we found that modern wheat breeding in China 419 significantly used the genetic diversity of the European landraces. In future breeding 420 programs, the genetic diversity of landraces from diverse geographic regions may 421 provide the additional genetic diversity required to fulfill the demands of our 422 increasing population and changing climate.

423 We found that different ancestors made distinct contributions to each 424 subgenome. In the Chinese cultivars, the genetic diversity of the A and B subgenomes 425 was mainly contributed by the European landraces. By contrast, the D subgenome of 426 the Chinese cultivars contains more diversity from local landraces. Although the 427 European landraces and the Chinese landraces have similar levels of genetic diversity 428 for each subgenome (Fig. 2d), the European landraces experienced continuous gene 429 exchange with the allotetraploid *T. turgidum*. Such a gene exchanges may diversify 430 favorable alleles that would be selected during modem breeding in China. On the 431 other hand, the D subgenome has limited genetic exchange in all landraces. As a 432 result, the D subgenome of the European landraces may not have additional 433 advantages over Chinese landraces as the A and B subgenomes have. These 434 differences may explain the different contribution of AB and D subgenomes to the 435 Chinese cultivars. Alternatively, the selection of accessions may lead to inevitable 436 bias, although we have used genome-wide markers to ensure that the resequenced 437 accessions represent the widest genome diversity.

We have narrowed down the selective sweeps corresponding to selection during wheat diversification and modern breeding, which will be useful for the future characterization of genes underlying important traits. Consistent with the contributions of different ancestors to each subgenome, we found that the

442 homoeologous genes in the different subgenomes are often under differential

selection, suggesting that it may be feasible to target a single copy of a homoeologousgene during breeding.

445

#### 446 Methods

#### 447 **Plant materials**

448 A total of 120 hexaploid wheat (*Triticum aestivum*) accessions from Asia, Europe,

449 and America were used in this study, including 95 landraces and 25 cultivars. Among

450 them were 60 accessions, including 42 landraces, and 18 elite varieties, from a

451 Chinese mini core collection, which is estimated to represent the majority of the

452 genomic diversity (~70%) of the 23,090 accessions in the Chinese national

453 collection<sup>16,56</sup>. Based on genome-wide simple sequence repeat (SSR) data, these 60

454 accessions were estimated to represent the widest genome diversity of the mini core

455 collection. The rest of the accessions were selected following the sequencing of 326

456 accessions collected from major wheat cultivation sites using the genotyping-by-

457 sequencing (GBS) method<sup>57</sup>. Based on the GBS results, the other 60 varieties were

458 selected for inclusion in the study, including 16 from West Asia, 12 from Central and

459 South Asia, 1 from Japan, 24 from Europe, and 7 from the Americas, which represent

460 the widest genome diversity. All the accessions were purified in multiple rounds of

461 single-seed descent to ensure their homozygosity.

462

#### 463 Library preparation for Illumina next-generation sequencing

464 Approximately 1.5 μg genomic DNA was extracted from each sample using the

465 CTAB method and prepared in libraries for sequencing using a TruSeq Nano DNA

466 HT sample preparation kit (Illumina, USA) following the manufacturer's

467 recommendations. Briefly, the genomic DNA samples were fragmented by sonication

468 to a size of ~350 bp, then end-polished, A-tailed, and ligated with the full-length

469 adapters for Illumina sequencing with further PCR amplification. Index codes were

470 added facilitate the differentiation of sequences from each sample. The PCR products

471 were purified (AMPure XP bead system; Beckman Coulter, USA) and the libraries

472 were analyzed for their size distribution using an Agilent 2100 Bioanalyzer (Agilent

- 473 Technologies, USA) and quantified using real-time PCR.
- 474

# 475 Genome sequencing and quality control

- 476 The libraries were sequenced using the Illumina HiSeq X platform (Illumina). In total,
- 477 ~21,676 Tb of raw data were generated. To ensure reliable reads without artificial
- 478 bias, the low-quality paired reads ( $\geq 10\%$  unidentified nucleotides (N); > 10
- 479 nucleotides aligned to the adaptor, allowing  $\leq 10\%$  mismatches; > 50% bases with a
- 480 Phred quality less than 5) were removed. A total of 21,531 Tb (~179.4 Gb per
- 481 sample) high-quality genomic data was obtained.
- 482

# 483 Read mapping and SNP calling

484 The remaining high-quality paired-end reads were mapped to the bread wheat 485 reference genome (IWGSC RefSeq v1.0) using Burrows-Wheeler Aligner software with the command 'mem -t 4 -k  $32 - M'^{58}$ . To reduce mismatches generated by the 486 487 PCR amplification before sequencing, the duplicated reads were removed with the help of SAMtools (v0.1.19)<sup>59</sup>. After the alignment, SNP calling was performed on a 488 489 population scale using a Bayesian approach in SAMtools. The genotype likelihoods 490 were calculated from the reads of each individual at each genomic location, and the allele frequencies were determined using a Bayesian approach. The 'mpileup' 491 492 command was used to identify SNPs using the parameters '-q 1 -C 50 -S -D -m 2 -F 493 0.002'. To exclude the SNP calling errors caused by incorrect mapping or InDels, 494 only the 70,172,660 high-quality filtered SNPs (depth  $\geq$  4, maf  $\geq$  0.01, miss  $\leq$  0.1) 495 were used in the subsequent analysis.

496

# 497 Functional annotation of genetic variants

The SNP annotation was performed using the published wheat genome<sup>9</sup> and the 498 ANNOVAR package (v2013-05-20)<sup>60</sup>. Based on the genome annotation, the SNPs 499 were categorized as being in exonic regions (overlapping with a coding exon), 500 501 intronic regions (overlapping with an intron), splicing sites (within 2 bp of a splicing 502 junction), upstream or downstream regions (within a 1-kb region upstream or 503 downstream of a transcription start or stop site, respectively), or intergenic regions. 504 The SNPs in the coding exons were further grouped into synonymous SNPs (do not cause amino acid changes) or non-synonymous SNPs (cause amino acid changes). 505 506 The mutations causing stop gain and stop loss were also classified into this group. To 507 exclude genes with possible structural annotation errors, those expressed in the leaves 508 or existing in multiple copies were selected as HC genes for further analysis. 509

- 510 **Population genetic diversity**
- 511 Nucleotide diversity  $\theta \pi$  and fixation index ( $F_{ST}$ ) were calculated using VCFtools

512  $(v0.1.14)^{61}$ , while Watterson's estimator ( $\theta_w$ ) and Tajima's D were calculated using

513 VariScan  $(v2.0.3)^{62}$ . These population statistics were analyzed using the sliding-

514 window approach (20-kb windows with 10-kb increments). HC genes were selected

to identify single-copy orthologous genes between the A, B, and D subgenomes using

516 Proteinortho (v5.16), with default settings $^{63}$ .

517

#### 518 Linkage disequilibrium analysis

519 To estimate and compare the patterns of linkage disequilibrium (LD) between

520 different populations, the squared correlation coefficient ( $r^2$ ) between pairwise SNPs

- 521 was computed using the software Haploview  $(v4.2)^{64}$ . The program parameters were
- 522 set as '-n -dprime -minMAF 0.1'. The average  $r^2$  value was calculated for pairwise
- 523 markers in a 500-kb window and averaged across the whole genome.
- 524

## 525 **Phylogenetic tree and population structure**

526 A total of 1,925,854 SNPs in coding regions (exonic and intronic) were used for the

527 population genetics analysis. To clarify the phylogenetic relationship from a genome-

- 528 wide perspective, an individual-based neighbor-joining tree was constructed using the
- 529 p-distance in the software TreeBeST (v1.9.2), with bootstrap values determined from
- 530 1000 replicates<sup>65</sup>.

531 The population genetic structure was examined using the program ADMIXTURE (v1.23)<sup>66</sup>. First, 95 landraces were used to estimate the genetic 532 533 ancestry, specifying a K ranging from 2 to 8. The most suitable number of ancestral populations was determined to be K = 3, for which the lowest cross-validation error of 534 535 0.492 was obtained. A principal component analysis (PCA) was also conducted to evaluate the genetic structure of the populations using the software GCTA<sup>67</sup>. First, a 536 537 genetic relationship matrix (GRM) was obtained using the parameter '-make-grm', then the top three principal components were estimated with the parameter '-pca3'. 538

539

#### 540 **Population admixture analyses**

The population relatedness and migration events were inferred using TreeMix<sup>68</sup>. A 541 542 total of 113 varieties with little admixture were selected to represent the seven 543 subgroups. The 1,925,854 coding-region SNPs were used to build a maximum 544 likelihood tree, using a window size of 2000 SNPs. This was repeated 10 times, using 545 the West Asian landraces as the root group. The tree with the lowest standard error for 546 the residuals was selected as the base tree topology. The population pairs with an 547 above zero standard residual error were identified as candidates for admixture events, 548 which represent populations which the data indicate are more closely related to each other than is demonstrated in the best-fit tree<sup>68</sup>. TreeMix was then run using between 549 550 one and six introduced migration events. When three migration events were added, 551 the residuals were much lower than for the trees generated using other numbers of 552 migration events.

553

## 554 Introgression analysis

The identity scores (IS) were calculated<sup>32</sup> to visualize the shared haplotypes between
the three populations (European varieties, Chinese landraces, and Chinese cultivars).

557 The IS values were used to evaluate the similarities of every sequenced sample to the 558 reference genome (Chinese Spring) within 20-kb windows. The IS was calculated

559 using the following formula:

560

561 where *D* represents genotype similarity to reference genome of samples in single site.

562 The IS of any single site was calculated as the difference in the D between two

samples. The average IS value was calculated for each population.

The  $H_P$  (population heterozygosity) was calculated for each candidate introgression region. At each detected SNP position, the numbers of reads corresponding to the most and least frequently observed allele (nMAJ and nMIN, respectively) were counted in each population. The  $H_P$  for each window was

568 calculated using the following formula:

569 
$$H_{\rm P} = \frac{2\sum n_{MAJ} \sum n_{MIN}}{(\sum n_{MAJ} + \sum n_{MIN})^2}$$

570  $\sum n_{MAJ}$  and  $\sum n_{MIN}$  are sums of the  $n_{MAJ}$  and  $n_{MIN}$  values, respectively, calculated 571 for all SNPs in the 20-kb window (sliding in 10-kb steps)<sup>69</sup>. The population 572 recombination rate was estimated from the SNPs of three populations (European 573 varieties, Chinese landraces, and Chinese cultivars) within a 20-kb window using the 574 R package FastEPRR (v1.0)<sup>70</sup>.

575

#### 576 Genome-wide selective sweep analysis

577 A sliding-window approach (20-kb windows sliding in 10-kb steps) was applied to

578 quantify the levels of polymorphism ( $\theta_{\pi}$ , the pairwise nucleotide variation as a

579 measure of variability) and genetic differentiation ( $F_{ST}$ ) between the different

- 580 populations using the VCFtools software (v0.1.14)<sup>61</sup>. The  $\theta_{\pi}$  ratios were log<sub>2</sub>-
- transformed. Subsequently, the empirical percentiles of the  $F_{ST}$  and  $\log_2(\theta_{\pi}$  ratio)
- 582 values in each window were estimated and ranked. The windows with the top 5%  $F_{ST}$
- and  $\log_2(\theta_{\pi} \text{ ratio})$  values were considered simultaneously as candidate outliers under
- 584 strong selective sweeps. All outlier windows were assigned to corresponding regions

and genes. The cross-population extended haplotype homozygosity (XP-EHH)

586 statistic was estimated<sup>71</sup> for the cultivar group and the landrace group, using the

587 landrace group as a contrast. The genetic map was assumed to be 0.18 cM/Mb for the

588 A and B subgenome, and 0.341 cM/Mb for the D subgenome. The XP-CLR score<sup>35</sup>

589 was used to confirm the selective sweeps on the basis of domestication features, with

the highest being 1%, via the cross-population composite likelihood method.

591

## 592 Candidate gene analysis

593 Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG)

analyses were performed on the candidate genes (all annotated genes in the outlier

595 windows denoted by the top 5% of the  $F_{ST}$  and  $\log_2(\theta_{\pi} \text{ ratio})$  values, the top 1% of the

596 XP-EHH values, and the top 1% of the XP-CLR scores). The candidate genes were

597 also attributed to known KEGG pathways (http://www.kegg.jp).

598

## 599 PCR primers and amplicon sequence analysis

600 A total of 28 Chinese cultivars developed between the 1940s and the 2000s were

analyzed to clarify the changes in the haplotypes of the candidate genes during wheat

breeding. All primers were designed using the Primer 5.0 software and listed in Table

603 S13a. DNAMAN8.0 was used to align the sequences to the reference genome and

604 identify the SNPs, after which the frequencies of the main haplotypes in the candidate

605 genes were analyzed.

606

## 607 Data availability

The sequencing data from this study have been deposited into the Sequence Read

609 Archive (https://www.ncbi.nlm.nih.gov/sra) under accession PRJNA439156.

610

# **References**

612	1.	Salamini, F., Ozkan, H., Brandolini, A., Schafer-Pregl, R. & Martin, W.
613		Genetics and geography of wild cereal domestication in the near east. Nat Rev
614		Genet <b>3</b> , 429-41 (2002).
615	2.	Gornicki, P. et al. The chloroplast view of the evolution of polyploid wheat.
616		New Phytol 204, 704-14 (2014).
617	3.	Dubcovsky, J. & Dvorak, J. Genome plasticity a key factor in the success of
618		polyploid wheat under domestication. Science 316, 1862-6 (2007).
619	4.	International Wheat Genome Sequencing Consortium. A chromosome-based
620		draft sequence of the hexaploid bread wheat (Triticum aestivum) genome.
621		Science <b>345</b> , 1251788 (2014).
622	5.	Marcussen, T. et al. Ancient hybridizations among the ancestral genomes of
623		bread wheat. Science 345, 1250092 (2014).
624	6.	Chapman, J.A. et al. A whole-genome shotgun approach for assembling and
625		anchoring the hexaploid bread wheat genome. Genome Biol 16, 26 (2015).
626	7.	Brenchley, R. et al. Analysis of the bread wheat genome using whole-genome
627		shotgun sequencing. Nature 491, 705-10 (2012).
628	8.	Clavijo, B.J. et al. An improved assembly and annotation of the allohexaploid
629		wheat genome identifies complete families of agronomic genes and provides
630		genomic evidence for chromosomal translocations. Genome Res 27, 885-896
631		(2017).
632	9.	International Wheat Genome Sequencing Consortium et al. Shifting the limits
633		in wheat research and breeding using a fully annotated reference genome.
634		Science <b>361</b> , eaar7191 (2018).
635	10.	Feldmann, M. Origin of cultivated wheat. in The world wheat book: a history
636		of wheat breeding., Vol. 1 (eds. Bonjean, A. & Angus, W.) 3-56 (Lavoisier
637		Publishing, Paris, France, 2001).
638	11.	Zhou, Y. et al. Uncovering the dispersion history, adaptive evolution and
639		selection of wheat in China. Plant Biotechnol J 16, 280-291 (2018).
640	12.	Long, T. et al. The early history of wheat in China from (14)C dating and
641		Bayesian chronological modelling. Nat Plants 4, 272-279 (2018).
642	13.	He, Z.H., Rajaram, S., Xin, Z.Y. & Huang, G.Z. A history of wheat breeding
643		in China, (CIMMYT, Mexico, D. F., 2001).
644	14.	Montenegro, J.D. et al. The pangenome of hexaploid bread wheat. Plant J 90,
645		1007-1013 (2017).
646	15.	Jordan, K.W. et al. A haplotype map of allohexaploid wheat reveals distinct
647		patterns of selection on homoeologous genomes. Genome Biol 16, 48 (2015).
648	16.	Hao, C. et al. Genetic diversity and construction of core collection in Chinese
649		wheat genetic resources. Chinese Sci Bull 53, 1518-1526 (2008).
650	17.	Li, H. A statistical framework for SNP calling, mutation discovery,
651		association mapping and population genetical parameter estimation from
652		sequencing data. Bioinformatics 27, 2987-93 (2011).

653	18.	Paterson, A.H. et al. The Sorghum bicolor genome and the diversification of
654		grasses. Nature 457, 551-6 (2009).
655	19.	McNally, K.L. et al. Genomewide SNP variation reveals relationships among
656		landraces and modern varieties of rice. Proc Natl Acad Sci USA 106, 12273-
657		8 (2009).
658	20.	Valliyodan, B. et al. Landscape of genomic diversity and trait discovery in
659		soybean. Sci Rep 6, 23598 (2016).
660	21.	Clark, R.M. et al. Common sequence polymorphisms shaping genetic
661		diversity in Arabidopsis thaliana. Science 317, 338-42 (2007).
662	22.	Akhunov, E.D. et al. Nucleotide diversity maps reveal variation in diversity
663		among wheat genomes and chromosomes. BMC Genomics 11, 702 (2010).
664	23.	Haudry, A. et al. Grinding up wheat: a massive loss of nucleotide diversity
665		since domestication. Mol Biol Evol 24, 1506-17 (2007).
666	24.	Gore, M.A. et al. A first-generation haplotype map of maize. Science 326,
667		1115-7 (2009).
668	25.	Xu, X. et al. Resequencing 50 accessions of cultivated and wild rice yields
669		markers for identifying agronomically important genes. <i>Nat Biotechnol</i> <b>30</b> ,
670		105-11 (2011).
671	26.	Mace, E.S. et al. Whole-genome sequencing reveals untapped genetic
672		potential in Africa's indigenous cereal crop sorghum. Nat Commun 4, 2320
673		(2013).
674	27.	Dvorak, J., Akhunov, E.D., Akhunov, A.R., Deal, K.R. & Luo, M.C.
675		Molecular characterization of a diagnostic DNA marker for domesticated
676		tetraploid wheat provides evidence for gene flow from wild tetraploid wheat to
677		hexaploid wheat. <i>Mol Biol Evol</i> <b>23</b> , 1386-1396 (2006).
678	28.	Zohary, D. & Brick, Z. <i>Triticum dicoccoides</i> in Israel: notes on its distribution,
679		ecology and natural hybridization. <i>Wheat Inform. Serv.</i> <b>13</b> , 6-8 (1961).
680	29.	Huang, X. <i>et al.</i> Genome-wide association studies of 14 agronomic traits in
681	_, .	rice landraces. <i>Nat Genet</i> <b>42</b> , 961-7 (2010).
682	30.	Hufford, M.B. <i>et al.</i> Comparative population genomics of maize
683	001	domestication and improvement. <i>Nat Genet</i> <b>44</b> , 808-11 (2012).
684	31.	Hwang, E.Y. <i>et al.</i> A genome-wide association study of seed protein and oil
685	511	content in soybean. <i>BMC Genomics</i> <b>15</b> , 1 (2014).
686	32.	Ai, H. <i>et al.</i> Adaptation and possible ancient interspecies introgression in pigs
687	52.	identified by whole-genome sequencing. <i>Nat Genet</i> <b>47</b> , 217-25 (2015).
688	33.	Devos, K.M., Dubcovsky, J., Dvorak, J., Chinoy, C.N. & Gale, M.D.
689	55.	Structural evolution of wheat chromosomes 4A, 5A, and 7B and its impact on
690		recombination. <i>Theor Appl Genet</i> <b>91</b> , 282-8 (1995).
690	34.	Sabeti, P.C. <i>et al.</i> Genome-wide detection and characterization of positive
691 692	54.	selection in human populations. <i>Nature</i> <b>449</b> , 913-8 (2007).
692	35.	Chen, H., Patterson, N. & Reich, D. Population differentiation as a test for
693 694	<i>JJ</i> .	selective sweeps. <i>Genome Res</i> <b>20</b> , 393-402 (2010).
074		serective sweeps. <i>Genome Res</i> <b>20</b> , <i>373</i> -402 (2010).

695	36.	Li, F. et al. Genome-wide linkage mapping of yield-related traits in three
696		Chinese bread wheat populations using high-density SNP markers. <i>Theor Appl</i>
697		Genet <b>131</b> , 1903-1924 (2018).
698	37.	Wang, Y.Y., Hsu, P.K. & Tsay, Y.F. Uptake, allocation and signaling of
699		nitrate. <i>Trends Plant Sci</i> <b>17</b> , 458-67 (2012).
700	38.	Buchner, P. & Hawkesford, M.J. Complex phylogeny and gene expression
701		patterns of members of the NITRATE TRANSPORTER 1/PEPTIDE
702		TRANSPORTER family (NPF) in wheat. J Exp Bot 65, 5697-710 (2014).
703	39.	Fang, Y. et al. A stress-responsive NAC transcription factor SNAC3 confers
704		heat and drought tolerance through modulation of reactive oxygen species in
705		rice. J Exp Bot 66, 6803-17 (2015).
706	40.	Gray, J.A., Shalit-Kaneh, A., Chu, D.N., Hsu, P.Y. & Harmer, S.L. The
707		<i>REVEILLE</i> clock genes inhibit growth of juvenile and adult plants by control
708		of cell size. <i>Plant Physiol</i> <b>173</b> , 2308-2322 (2017).
709	41.	Peterson, K.M. et al. Arabidopsis homeodomain-leucine zipper IV proteins
710		promote stomatal development and ectopically induce stomata beyond the
711		epidermis. Development 140, 1924-35 (2013).
712	42.	Campbell, J.L. et al. Cloning of new members of heat shock protein HSP101
713		gene family in wheat (Triticum aestivum (L.) Moench) inducible by heat,
714		dehydration, and ABA. Biochim Biophys Acta 1517, 270-7 (2001).
715	43.	Hu, H. et al. Characterization of transcription factor gene SNAC2 conferring
716		cold and salt tolerance in rice. Plant Mol Biol 67, 169-81 (2008).
717	44.	Sun, T., Li, S. & Ren, H. OsFH15, a class I formin, interacts with
718		microfilaments and microtubules to regulate grain size via affecting cell
719		expansion in rice. Sci Rep 7, 6538 (2017).
720	45.	Ferrari, S., Galletti, R., Vairo, D., Cervone, F. & De Lorenzo, G. Antisense
721		expression of the Arabidopsis thaliana AtPGIP1 gene reduces
722		polygalacturonase-inhibiting protein accumulation and enhances susceptibility
723		to Botrytis cinerea. Mol Plant Microbe Interact 19, 931-6 (2006).
724	46.	Mukhtar, M.S., Deslandes, L., Auriac, M.C., Marco, Y. & Somssich, I.E. The
725		Arabidopsis transcription factor WRKY27 influences wilt disease symptom
726		development caused by Ralstonia solanacearum. Plant J 56, 935-47 (2008).
727	47.	Xu, P. et al. A brassinosteroid-signaling kinase interacts with multiple
728		receptor-like kinases in Arabidopsis. Mol Plant 7, 441-4 (2014).
729	48.	Piechota, J. et al. Unraveling the functions of type II-prohibitins in
730		Arabidopsis mitochondria. Plant Mol Biol 88, 249-67 (2015).
731	49.	Talboys, P.J., Healey, J.R., Withers, P.J. & Jones, D.L. Phosphate depletion
732		modulates auxin transport in Triticum aestivum leading to altered root
733		branching. J Exp Bot 65, 5023-32 (2014).
734	50.	Liu, G. et al. Profiling of wheat class III peroxidase genes derived from
735		powdery mildew-attacked epidermis reveals distinct sequence-associated
736		expression patterns. Mol Plant Microbe Interact 18, 730-41 (2005).

737	51.	Lori, M. et al. Evolutionary divergence of the plant elicitor peptides (Peps)
738		and their receptors: interfamily incompatibility of perception but compatibility
739		of downstream signalling. J Exp Bot 66, 5315-25 (2015).
740	52.	Jin, H. et al. Genome-wide QTL mapping for wheat processing quality
741		parameters in a Gaocheng 8901/Zhoumai 16 recombinant inbred line
742		population. Front Plant Sci 7, 1032 (2016).
743	53.	Calderón-Villalobos, L.I., Nill, C., Marrocco, K., Kretsch, T. &
744		Schwechheimer, C. The evolutionarily conserved Arabidopsis thaliana F-box
745		protein AtFBP7 is required for efficient translation during temperature stress.
746		<i>Gene</i> <b>392</b> , 106-16 (2007).
747	54.	Savatin, D.V. et al. The Arabidopsis NUCLEUS- AND PHRAGMOPLAST-
748		LOCALIZED KINASE1-related protein kinases are required for elicitor-
749		induced oxidative burst and immunity. Plant Physiol 165, 1188-1202 (2014).
750	55.	Morimoto, R., Nishioka, E., Murai, K. & Takumi, S. Functional conservation
751		of wheat orthologs of maize rough sheath1 and rough sheath2 genes. Plant
752		Mol Biol 69, 273-85 (2009).
753	56.	Hao, C., Wang, L., Ge, H., Dong, Y. & Zhang, X. Genetic diversity and
754		linkage disequilibrium in Chinese bread wheat (Triticum aestivum L.) revealed
755		by SSR markers. PLoS ONE 6, e17279 (2011).
756	57.	Poland, J.A., Brown, P.J., Sorrells, M.E. & Jannink, J.L. Development of
757		high-density genetic maps for barley and wheat using a novel two-enzyme
758		genotyping-by-sequencing approach. PLoS ONE 7, e32253 (2012).
759	58.	Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-
760		Wheeler transform. <i>Bioinformatics</i> 25, 1754-60 (2009).
761	59.	Li, H. et al. The Sequence Alignment/Map format and SAMtools.
762		<i>Bioinformatics</i> 25, 2078-9 (2009).
763	60.	Wang, K., Li, M. & Hakonarson, H. ANNOVAR: functional annotation of
764		genetic variants from high-throughput sequencing data. Nucleic Acids Res 38,
765		e164 (2010).
766	61.	Danecek, P. et al. The variant call format and VCFtools. Bioinformatics 27,
767		2156-8 (2011).
768	62.	Vilella, A.J., Blanco-Garcia, A., Hutter, S. & Rozas, J. VariScan: Analysis of
769		evolutionary patterns from large-scale DNA sequence polymorphism data.
770		<i>Bioinformatics</i> <b>21</b> , 2791-3 (2005).
771	63.	Lechner, M. <i>et al.</i> Proteinortho: detection of (co-)orthologs in large-scale
772		analysis. <i>BMC Bioinformatics</i> <b>12</b> , 124 (2011).
773	64.	Barrett, J.C., Fry, B., Maller, J. & Daly, M.J. Haploview: analysis and
774	~ =	visualization of LD and haplotype maps. <i>Bioinformatics</i> <b>21</b> , 263-5 (2005).
775	65.	Vilella, A.J. <i>et al.</i> EnsemblCompara GeneTrees: Complete, duplication-aware
776		phylogenetic trees in vertebrates. <i>Genome Res</i> <b>19</b> , 327-35 (2009).
777 779	66.	Alexander, D.H., Novembre, J. & Lange, K. Fast model-based estimation of
778		ancestry in unrelated individuals. Genome Res 19, 1655-64 (2009).

779	67.	Yang, J., Lee, S.H., Goddard, M.E. & Visscher, P.M. GCTA: a tool for
780		genome-wide complex trait analysis. Am J Hum Genet 88, 76-82 (2011).
781	68.	Pickrell, J.K. & Pritchard, J.K. Inference of population splits and mixtures
782		from genome-wide allele frequency data. PLoS Genet 8, e1002967 (2012).
783	69.	Rubin, C.J. et al. Whole-genome resequencing reveals loci under selection
784		during chicken domestication. Nature 464, 587-91 (2010).
785	70.	Gao, F., Ming, C., Hu, W. & Li, H. New software for the fast estimation of
786		population recombination rates (FastEPRR) in the genomic era. G3 6, 1563-71
787		(2016).
788	71.	Sabeti, P.C. et al. Genome-wide detection and characterization of positive
789		selection in human populations. Nature 449, 913-918 (2007).
790		

# 791 Acknowledgments

792 We thank Yalong Guo and Wenfeng Qian for their valuable suggestions. This

research was supported by the Chinese Academy of Sciences (CAS; grant no.

794 XDA08020105 to Y.J. and XDA08040108 to H.C.), the National Natural Science

Foundation of China (grant no. 31430010 to Y.J., 31871245 to Ying W, and

796 31770311 to C.T.), the National Program for Support of Top-Notch Young

797 Professionals (Y.J.), University of CAS (grant no. 110601M206 to Ying W.), the

798 Beijing NOVA Program (grant no. Z161100004916107 to Ying W.), the National

799 Transgenic Science and Technology Program (grant no. 2016ZX08010-002 to C.T.),

and the CAS Youth Innovation Promotion Association (grant no. 2017139 to C.T.).

801

# 802 Author information

803 These authors contributed equally: H. Chen, C. Jiao, Ying Wang, Yuange Wang.

804

# 805 Affiliantions

806 State Key Laboratory of Plant Genomics, Institute of Genetics and Developmental

- 807 Biology, Chinese Academy of Sciences, Beijing 100101, China
- 808 Haofeng Chen, Yuange Wang, Caihuan Tian, Haopeng Yu, Jing Wang, Yuling Jiao

- 809 Novogene Bioinformatics Institute, Beijing 100083, China
- 810 Chengzhi Jiao, Wenkai Jiang, Hongfeng Lu
- 811 College of Life Sciences, University of Chinese Academy of Sciences, Beijing 100049,
- 812 China
- 813 Ying Wang, Fei Lu, Xiangdong Fu, Yongbiao Xue, Hongqing Ling, Yuling Jiao
- 814 West China Biomedical Big Data Center, West China Hospital/West China School of
- 815 Medicine, and Medical Big Data Center, Sichuan University, Chengdu 610041, China
- 816 Haopeng Yu
- 817 Department of Crop Genomics and Bioinformatics, College of Agronomy and
- 818 Biotechnology, National Maize Improvement Center of China, China Agricultural
- 819 University, Beijing 100193, China
- 820 Xiangfeng Wang
- 821 State Key Laboratory of Plant Cell and Chromosome Engineering, Institute of
- 822 Genetics and Developmental Biology, Chinese Academy of Sciences, Beijing 100101,
- 823 China
- Fei Lu, Xiangdong Fu, Yongbiao Xue, Hongqing Ling
- 825

## 826 **Contributions**

- 827 Y.J., H.Lu and Ying W. conceived of and designed the study. C.J., Yuange W., H.Y.,
- 828 J.W., X.W., W.J., and H.Lu performed the analyses. C.J., H.Lu, Y.J., Ying W., W.J.,
- 829 and H.Ling interpreted the data. H.C. Yuange W., C.T., J.W. F.L., X.F., Y.X., H.Ling,
- and Y.J. contributed to data collection. Y.J., C.J., and H.Lu wrote the manuscript with
- 831 input from all coauthors.

832

#### 833 Competing interests

834 The authors declare no competing interests.

bioRxiv preprint doi: https://doi.org/10.1101/519587; this version posted January 14, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

# 835

# 836 Corresponding authors

- 837 Correspondence to Yuling Jiao (<u>yljiao@genetics.ac.cn</u>) or Hongfeng Lu
- 838 (<u>luhongfeng@novogene.cn</u>).

# 840 **Table**

**Table 1.** Summary of the SNP distribution in the A, B, and D subgenomes.

	Up- stream <sup>a</sup>	Exonic									5'				
Sub- genome		Stop gain	Stop loss	Syno- nymous	Non-sy- nonymous	N/S ratio	Un- knowns	Intronic	3' UTR	5' UTR	UTR; 3' UTR <sup>b</sup>	Splicing	Down- stream <sup>c</sup>	Up/down Stream <sup>d</sup>	Intergenic
А	325,784	8,180	1,782	133,267	207,728	1.56	27,325	366,052	11,108	12,388	1	1,466	302,952	46,595	27,276,394
В	349,547	9,130	1,953	141,373	224,779	1.59	30,931	394,269	11,557	14,315	11	1,615	324,819	52,059	31,395,654
D	124,077	3,487	644	82,817	105,693	1.28	8,690	151,873	5,590	5,881	4	660	117,057	18,823	7,287,737
Unknown	11,337	327	58	6,529	9,168	1.40	838	8,961	446	299	0	44	10,984	1,397	536,205
Total	810,745	21,124	4,437	363,986	547,368	1.50	67,784	921,155	28,701	32,883	16	3,785	755,812	118,874	66,495,990

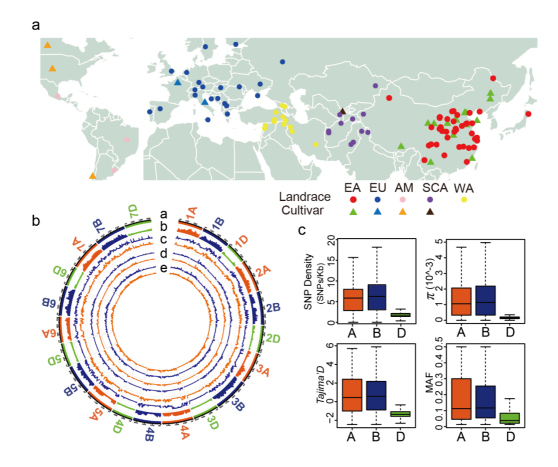
<sup>a</sup>Regions within the 1 kb before the start codon of the downstream gene.

<sup>b</sup>Regions considered as both the 3'UTR of the upstream gene and 5'UTR of the downstream gene.

844 <sup>c</sup>Regions within the 1 kb after the stop codon of the upstream gene.

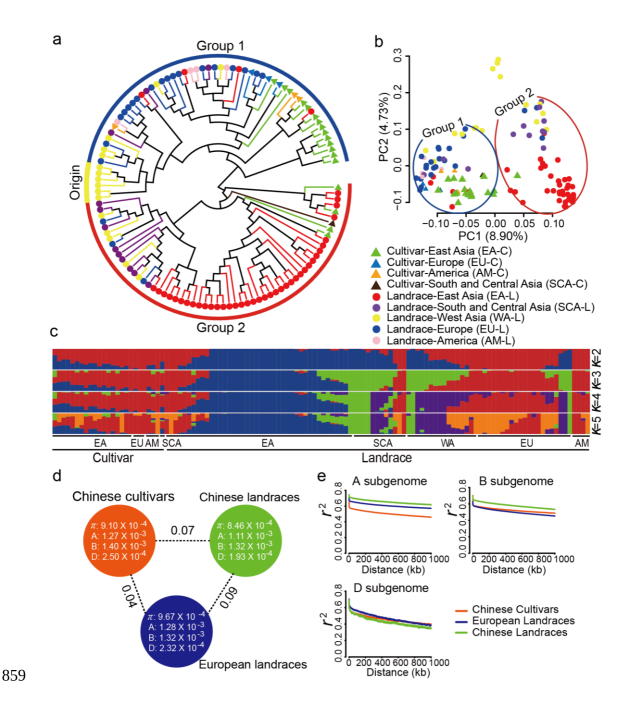
<sup>d</sup>Regions within the 1 kb after the stop codon of the upstream gene and also within the 1 kb before the start codon of the downstream gene.

# 846 Figures



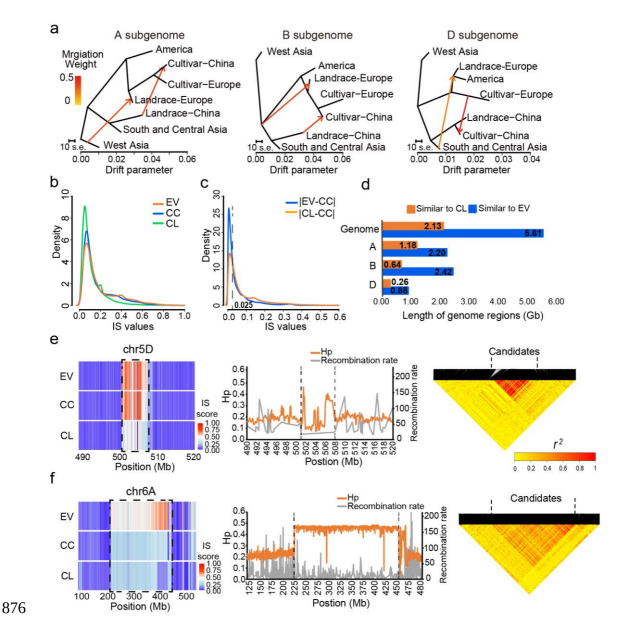
847

848 Fig. 1. Characterization of SNP distribution across the A, B, and D subgenome 849 chromosomes. (a) The geographic distribution of the 120 wheat accessions used in 850 this study. The circles represent landraces, while triangles represent cultivars. 851 Different colors represent populations from different geographic regions. EA, East 852 Asia; WA, West Asia; EU, Europe; AM, America; SCA, South and Central Asia. (b) 853 The genetic diversity of different populations along the chromosomes. From a to e, 854 the sections respectively represent the chromosomes, SNP density, neutral evolutionary parameters (Tajima's D), nucleotide diversity parameters ( $\theta_{\pi}$ ), and 855 856 Watterson's  $\theta$ , respectively. The red and blue lines in sections c to e represent the 857 cultivars and landraces, respectively. (c) Comparison of the genomic characteristics of the A, B, and D subgenomes. MAF, minor allele frequencies. 858



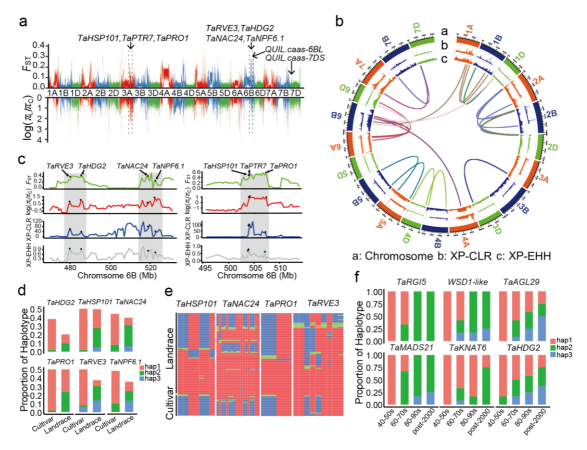
860 Fig. 2. Population structure of the 120 wheat accessions. (a) Phylogenetic tree of the wheat accessions used in this study. The triangles represent wheat cultivars, while 861 circles represent the landraces. The different colors represent populations from 862 different geographic locations. "Origin" represents West Asia, which is considered the 863 864 origin of bread wheat. "Group 1" represents the genotypes derived from the westward 865 migration of bread wheat, including the European and American landraces, and the cultivars from Europe, America, and China. "Group 2" represents the groups derived 866 867 from the eastward migration of bread wheat, including the South, Central, and East

- 868 Asian landraces. (b) Principal component analysis of the 120 wheat accessions. (a)
- and (**b**) share the same key. (**c**) Population structure of the 120 wheat accessions. (**d**)
- 870 Nucleotide diversity ( $\pi$ ) and population divergence ( $F_{ST}$ ) across the Chinese cultivars,
- 871 Chinese landraces, and European landraces. The values in each circle represent a
- 872 measure of nucleotide diversity for this group and each of its subgenomes, while the
- values on each line indicate the population divergences between the two compared
- groups. (e) Decay of linkage disequilibrium (LD) in the A, B, and D subgenomes of
- three populations.



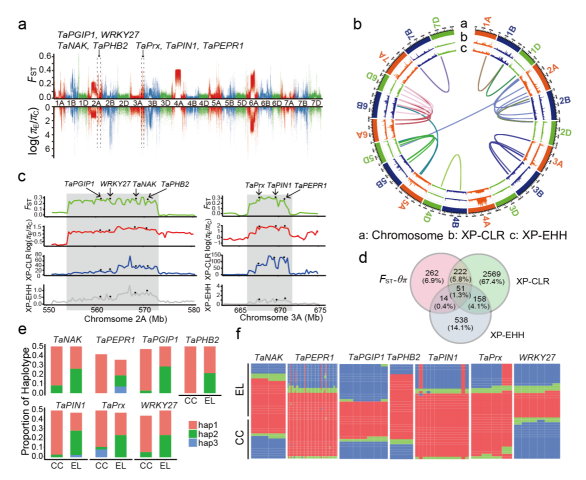
877 Fig. 3. Haplotype introgressions from the European and Chinese landraces into the Chinese cultivars. (a) Inferred phylogenetic trees of the A, B, and D subgenomes of 878 879 different populations, including mixture events. Horizontal branch lengths are 880 proportional to the amount of genetic drift. Migration arrows are colored according to 881 their weight. (b) Density distribution of identity score (IS) values for three 882 populations. (c) Density distribution of the difference in IS values between two 883 populations. |EL-CC| represents the absolute value of the IS differences between the European varieties and the Chinese cultivars. |CL-CC| represents the absolute value of 884 885 the IS differences between the Chinese landraces and the Chinese cultivars. (d)

886 Amount of similar genomic regions between populations. Orange represents the 887 length of similar genomic regions between the Chinese cultivars and the Chinese 888 landraces, and blue represents the length of similar genomic regions between the 889 Chinese cultivars and the European varieties. The statistical results of the entire 890 genome (Genome) as well as the individual A, B, and D subgenomes. The numbers 891 represent the total length of the similar genomic regions (Gb). (e) Candidate 892 introgression region between the European varieties and the Chinese cultivars. The left panel presents details of the 8-Mb candidate region, located on chromosome 5D, 893 894 in which the Chinese cultivars are more similar to the European varieties than the 895 Chinese landraces. The middle panel represents the distribution of the population 896 heterozygosity  $(H_P)$  and recombination rates in this candidate region. The right panel 897 represents the LD heatmap of this candidate region. (f) Candidate introgression region 898 (200 Mb, located on chromosome 6A) between the Chinese landraces and Chinese 899 cultivars. The central and right panels are as described for (e).



900

901 Fig. 4. Population sweep selection during modern breeding between the cultivars and 902 landraces. (a) The distribution of the  $F_{\rm ST}$  and  $\theta_{\pi}$  ratio (landrace/cultivar) along the 903 chromosomes. The genome-wide threshold was defined by the top 5% of the  $F_{ST}$  and  $\theta_{\pi}$  ratio (landrace/cultivar) values. Candidate genes and QTLs were marked on the 904 map. (b) The distribution of XP-CLR and XP-EHH scores along the chromosomes. 905 906 From the outside to the inside, the data presented are the XP-CLR score, XP-EHH 907 score, and the duplicated copies of the homoeologous genes in the different 908 subgenomes under selection for every chromosome. (c) Seven genes distributed in 909 three candidate regions were commonly found to be under selection using the  $F_{\rm ST}$ ,  $\theta_{\pi}$ 910 ratio, XP-CLR, and XP-EHH methods. The black points indicate the location of the 911 candidate selected genes on the chromosomes. (d) The haplotype frequency of the 912 candidate genes between the Chinese landrace and cultivar populations. (e) Heatmap of genotypes in the putative selective sweeps. (f) The selection of the favored 913 914 haplotype frequencies in modern Chinese cultivars during wheat breeding since the 915 1940s.



917 Fig. 5. Population sweep selection during local adaption in Chinese cultivars and European landraces. (a) The distribution of the  $F_{ST}$  and  $\theta_{\pi}$  ratio (European 918 919 landrace/Chinese cultivar) along the chromosomes. The genome-wide threshold was 920 defined by the top 5% of the  $F_{ST}$  and  $\theta_{\pi}$  ratio (European landrace/Chinese cultivar) 921 values. Candidate genes and QTLs are marked on the map. (b) The distribution of 922 XP-CLR and XP-EHH scores along the chromosomes. From the outside to the inside, 923 the data presented are the XP-CLR score, the XP-EHH score, and the duplicated 924 copies of the homoeologous genes in the different subgenomes under selection along 925 every chromosome. (c) Seven genes distributed in the two candidate selection regions 926 identified using the  $F_{\text{ST}}$ - $\theta_{\pi}$  ratio, XP-CLR, and XP-EHH methods. The black points 927 indicate the location of the candidate genes on the chromosome; every gene has a 928 strong selection signal. (d) A Venn diagram of wheat genes under selection detected 929 using the  $F_{\rm ST}$ - $\theta_{\pi}$  ratio, XP-CLR, and XP-EHH methods. (e) The haplotype frequency

bioRxiv preprint doi: https://doi.org/10.1101/519587; this version posted January 14, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

- 930 of the candidate genes between the Chinese cultivars (CC) and the European
- 931 landraces (EL). (f) Heatmap of genotypes in the putative selective sweep.