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Genomic analyses provide insights into peach local

2 adaptation and responses to climate change

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- 28 **Running title:** Genetic bases of peach local adaptation
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- 30 **Key words:** Peach, Whole-genome selection scan, Genome-wide environmental association study,
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34 The environment has constantly shaped plant genomes, but the genetic bases underlying 35 how plants adapt to environmental influences remain largely unknown. We constructed a 36 high-density genomic variation map by re-sequencing genomes of 263 geographically 37 representative peach landraces and wild relatives. A combination of whole-genome 38 selection scans and genome-wide environmental association studies (GWEAS) was 39 performed to reveal the genomic bases of peach local adaptation to diverse climates 40 comprehensively. A total of 2,092 selective sweeps that underlie local adaptation to both 41 mild and extreme climates were identified, including 339 sweeps conferring genomic 42 pattern of adaptation to high altitudes. Using GWEAS, a total of 3,496 genomic loci strongly 43 associated with 51 specific environmental variables were detected. The molecular 44 mechanism underlying adaptive evolution of high drought, strong UV-B, cold hardiness, 45 sugar content, flesh color, and bloom date were revealed. Finally, based on 30 years of 46 observation, a candidate gene associated with bloom date advance, representing peach 47 responses to global warming, was identified. Collectively, our study provides insights into 48 molecular bases of how environments have shaped peach genomes by natural selection 49 and adds valuable genome resources and candidate genes for future studies on 50 evolutionary genetics, adaptation to climate changes, and future breeding.

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52 Environmental adaptation is fundamental to species survival and conservation of biodiversity. 53 especially under threats of climate change (Blanquart et al. 2013). Unlike animals, which can 54 escape from hostile environments, plants are sessile and have to adapt by shaping and/or fixing 55 genetic variants that are conducive for survival. Generally, climate is the major selective pressure 56 driving adaptive evolution, resulting in different ecotypes within a single species (Hancock et al. 57 2011; Fournier-Level et al. 2011). However, the mechanisms underlying how climate shapes plant 58 genomes remain largely unclear. Recently, identifying adaptive variants and understanding 59 molecular mechanism of adaptation across a genome have become tractable due to the advances 60 of sequencing technologies. Recent studies have sought to elucidate genetic bases of adaptation 61 through genome-wide identification of regions under positive selection and/or loci that control 62 adaptive traits in Arabidopsis thaliana (Fournier-Level et al. 2011), rice (Yan et al. 2013), sorghum 63 (Lasky et al. 2015), and poplar (Wang et al. 2018). However, no study has focused on genetic 64 bases of adaptation in domesticated perennial fruit crops. Domesticated crops have adapted to 65 diverse climates during domestication and subsequent spread, and show local adaptation through 66 long-term natural selection. Landraces and wild relatives harbor great genetic diversity and an 67 abundance of resistance genes, which provide excellent resources for breeding initiatives. This is

especially the case with accessions originating from stressful environments that may have specific stress-resistance genes (Bolger et al. 2014a). However, a cost of domestication is that many resistance related genes have been lost. In addition, global climate change is driving decreases in productivity and distribution changes in several crop species (Tim and Braun, 2013). Therefore, it is of great importance to identify adaptive genes that can contribute to crop improvement, species survival, and global food security in the face of environmental deterioration.

74Peach is an important temperate fruit species, with a global yield of 24.7 million tons in 75 2017 (FAOSTAT; http://www.fao.org/faostat). It is also an important model system for the 76 Rosaceae family, members of which provide one of world's main resources of fruits. Peach 77 originated in southwestern China, and its landraces and wild relatives are widespread in both 78 temperate and sub-tropical regions, as well as in wet and dry climates (Wang et al. 2012). On the 79 grounds of wide distributions, peach can be regarded as an excellent material for studying 80 adaptation genetics. Peach has a relatively small genome size (~227.4 Mb) (Verde et al. 2013) 81 and genomic analyses have identified a number of loci and candidate genes associated with 82 human selection and agronomically important traits (Cao et al. 2014; Cao et al. 2016; Li et al. 83 2019). However, there have been few studies describing genomic loci associated with 84 environmental adaptation and natural selection.

To investigate the genetic basis of local adaptation, we sequenced a wide collection of 263 peach accessions from a broad range of geographical origins and associated with diverse climates, spanning mild and extreme environments. Using the sequencing data, we deciphered adaptive patterns across peach genome by combining the identification of signatures of selective sweeps with genome-wide association studies of environmental variables and adaptive traits. Finally, we identified a candidate gene associated with peach responses to global warming, based on observations over a 30-year period.

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93 **Results and discussion**

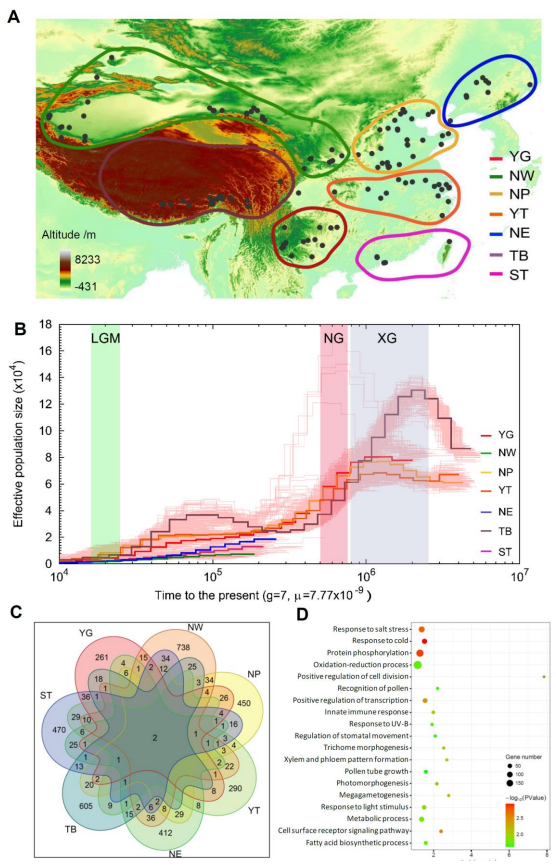
94 Genomic variation map and population structure

We first constructed a genome variation map for peach using a collection of 263 diverse accessions (Fig. 1A), consisting of 52 wild relatives and 211 landraces (Supplementary Table S1), which collectively capture more than 95% of geographic diversity of native distribution of peach landrace and wild relatives. A total of 342.7 Gb of sequence was generated, with a median depth of 5.3 × and coverage of 91.7% of reference peach 'Lovell' genome (release v2.0) (Verde et al. 2013) (Supplementary Table S1). We identified a final set of 4,611,842 high-quality single101 nucleotide polymorphisms (SNPs) (Supplementary Fig. S1A), of which 1,931,310 were intronic 102 (~11.33%) and 848,638 (~4.98%) were exonic. Among SNPs in coding sequence, we found that 103 7.853 SNPs present in 5.512 peach genes (~20.5% of total genes) are likely to have a major 104 impact on gene function. The accuracy of identified SNPs was found to be ~95.6%, based on 105 genotyping of 18 randomly selected SNPs in 130 accessions using a Sequenom MassARRAY 106 platform (Supplementary Table S2). In addition, we identified 1,049,266 small insertions and 107 deletions (INDELs) (shorter than or equal to 6 bp) and 106,388 large structural variations (SVs) 108 (> 30 bp) (Supplementary Fig. S1A).

109 We explored the genetic relationships among 263 accessions using 2,468,307 SNPs with 110 minor allele frequency (MAF) greater than 0.05. Based on the neighbor-joining tree and population 111 structure analysis, the 263 peach accessions could be divided into seven major groups, which 112 were largely congruent with ecotypes classified according to their geographic information, 113 including YG (Yun-gui Plateau), NW (Northwest China), NP (North Plain China), YT (Yangtze 114 River Middle and Backward), NE (Northeast China), TB (Tibet plateau), and ST (South China 115Sub-tropical) groups (Supplementary Fig. S1B, Supplementary Fig. S2, and Supplementary Table 116 S1). Although the neighbor-joining tree largely supported the division of seven major groups, there 117 were some discrepancies between geographical characterization and phylogenetic clustering 118 (Supplementary Fig. S2D), indicating shared ancestral variation and historical gene flow among 119 landraces in closely related groups. Moreover, principal component analysis (PCA) and model-120 based clustering analyses also supported the extensive admixture and possible gene flow among 121 landrace groups (Supplementary Fig. S2E and S2F). Furthermore, we found the small pair-wise 122 genetic differentiation (F_{ST}) values between different landrace groups, again consistent with 123 population admixture (Supplementary Fig. S2G).

Using the demographic analysis with the pairwise sequential Markovian coalescent (PSMC) model (Li and Durbin 2011), we found the sharply decline of effective population size (*Ne*) during the two largest Pleistocene glaciations: the Xixiabangma glaciation (1.17-0.8 MYA) and Naynayxungla glaciation (0.78-0.50 MYA), and a slight decline of *Ne* during the last glacial maximum (~20,000 years ago) (Fig. 1B).

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136 Fig. 1 Summary of 263 samples and genes under selection for seven peach groups. (A) 137 Geographic distribution of 263 peach accessions used in this study. Each accession is represented 138 by a dot on the world map. Seven ecotypes are highlighted using rings with different colors. (B) 139 Demographic history of the seven peach groups. Ancestral population size was inferred using the 140 PSMC model. Three periods of the last glacial maximum (LGM, ~20 KYA). Navnavxungla Glaciation 141 (NG, 0.5~0.78 MYA), and Xixiabangma Glaciation (XG, 0.8~0.17 MYA) are shaded in green, red, and 142 blue, respectively. (C) Venn diagram showing the number of genes under selection in the seven 143 groups. (D) Over-represented gene ontology (GO) terms in overall selection regions. Only the top 20 144 most over-represented terms are shown. YG, Yun-gui Plateau. NW, Northwest China. NP, North Plain 145 China. YT, Yangtze River Middle and Backward. NE, Northeast China. TB, Tibet plateau. ST, South 146 China Sub-tropical.

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148 Selective sweeps related to adaptation to diverse environments

149 Peach accessions of each group have adapted locally through long-term selection under local 150 environments (Supplementary Table S3). To identify genomic loci that favor local adaptation for 151seven groups, we detected signatures of selective sweeps for each group. This revealed a total 152of 2.092 genomic regions (19.1 Mb, ~8.4%; 189, 387, 301, 235, 280, 339, and 378 regions for the 153YG, NW, NP, YT, NE, TB, and ST groups, respectively) (Supplementary Fig. S3), which were 154 termed candidate selection regions (CSRs) (Supplementary Table S4). The overall CSRs 155harbored 4,198 genes (~17.5%), including 506, 1,197, 835, 530, 747, 920, and 869 genes for the 156 YG, NW, NP, YT, NE, TB, and ST groups, respectively (Fig. 1C). Selections on these genes may 157underlie the adaptation to different climates. Notably, we found that few genes were shared among 158different groups (Fig. 1C), suggesting the unique adaptive patterns for each group and that 159different climates may shape distinct genomic regions.

160 We found that genes related to response to different types of stimuli and stress, including 161 temperature, radiation, salt, DNA damage, osmotic, toxin, were overrepresented (P < 0.05), 162 suggesting that stress-related genes have participated in adaptive evolution (Fig. 1D, 163 Supplementary Table S5). For instance, two cation/H⁺ exchanger family genes (CHX) 164 (Prupe.6G251600 and Prupe.6G251700) and one salt overly sensitive 3 (SOS3) 165 (*Prupe.2G188700*) gene showed high reduction of diversity (ROD) and F_{ST} values in the NW 166 group. Homologs of these genes are involved in salt resistance in A. thaliana (Monihan et al. 167 2016), suggesting their potential contributions to adaptation to saline soils in northwestern China. 168 The resistance-related LRR (leucine-rich repeat) domain and PPR (pentatricopeptide repeat) 169 gene family were highly enriched in CSRs (P < 0.05) (Supplementary Table S5). The LRR domain,

which is considered to be one of the most important domains for plant resistance genes, was also enriched (P < 0.05), with 121 of 612 members (19.8%) in CSRs. PPR proteins form one of the largest protein families in land plants that are related to environmental responses, with 286 members in peach genome, of which 79 (~27.6%) were in CSRs.

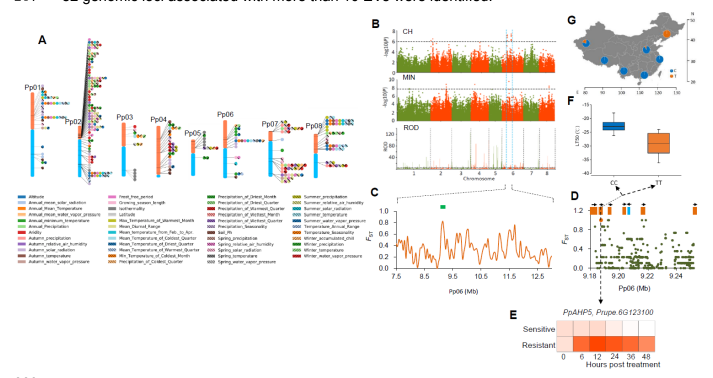
174The known genes or biological pathways involved in adaptation to the environment in the 175habitat of each group were determined. For instance, the YG group was distributed on the Yun-176 gui plateau (Southwest China), a low-latitude and high-altitude (~2000 m) region with high annual 177precipitation (> 1100 mm) and acidic soil (pH 4.5~5.5) (Supplementary Table S3). Genes related 178to metal ion (including potassium, iron, and zinc) binding and transport, cell membrane function, 179 and response to toxins were overrepresented in this group (107 genes, P < 0.05) (Supplementary 180 Table S5), consistent with functions in overcoming cation deficiency and aluminum toxicity that 181 are common in acidic soils (Seguel et al. 2013). For the YT group, we observed enrichments of 182 the LRR domain (24 genes), NB-ARC domain (8 genes), and other genes related to stress 183 responses (32 genes) (P < 0.05) (Supplementary Table S5), in comparison to other groups. This 184 suggests that the YT group has accumulated more abiotic and biotic stress-resistance variants 185 due to strong selective pressures in high temperature and high humidity areas (Supplementary 186 Table S3). These results indicate that accessions from the YT group may exhibit higher 187 adaptability than other landrace accessions.

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189 Genome-wide environmental association studies of 51 environmental variables

190 Although we obtained candidate genes underlying adaptation by identifying selective sweeps, 191 many adaptive events in natural populations may occur by polygenic adaptation, which would be 192 largely undetected by conventional methods for detecting selection (Pritchard and Di Rienzo 193 2010). However, local adaptation can generate correlations between environmental variables 194 (EVs) and genomic loci which can be used to detect polygenic adaptation. We investigated a total 195 of 51 EVs of the geographic origin of each accession that are important for plant adaptation 196 (Supplementary Table S6 and S7), representing extremes and seasonality of temperature and 197 precipitation, latitude, altitude, relative air humidity, water vapor pressure, growing season lengths, 198 and radiations. Using a Mantel test, we found a significant correlation between geographic and 199 genetic distances (Pearson's r = 0.73, P = 0.000999), with most associations being driven by 200 altitude. To obtain loci associated with EVs, we performed GWEAS on 51 EVs. A total of 9393 201 association SNPs (Supplementary Table S8), involving 3807 genes, were identified (Fig. 2A). 202 Notably, we found an EV association hotspot regions at the top of chromosome 2 that was

enriched with genes encoding NB-LRR proteins in peach genome (Verde et al. 2013). Consistent
 with the high correlations among some climate variables (Supplementary Fig. S4), only 3496
 association SNPs were unique, and ~62.8% of the associations were shared across different
 types of EVs, suggesting that different EVs may shape same genomic regions. Notably, a total of
 82 genomic loci associated with more than 10 EVs were identified.



223 Fig. 2 Genome-wide environmental association studies of 51 environmental variables and 224 genomic loci associated with winter cold adaptation. (A) SNPs associated with environmental 225 variables (EVs). Only the top 10 association signals for each EV are shown. All signals were included 226 if the total number of signals was < 10. (B) The *PpAHP* locus involved in adaptation to winter low 227 temperature in peach. Manhattan plots for a GWAS study of cold hardiness (CH) and winter lowest 228 temperature (MIN), and selection signals of the NE group (ROD) were detailed. The dashed 229 line represents the significance threshold for each test. The candidate genomic region is highlighted 230 between two dashed blue vertical lines. (C) Distribution of F_{ST} values between NE and ST groups in 231 the candidate region. The green bar indicates the *PpAHP* locus. (**D**) Close-up view of the F_{ST} values 232 in a region corresponding to the green bar in (C). This region contains six *PpAHP* homologs (orange) 233 and one other gene (light blue). The candidate SNP is highlighted using an orange dot. (E) Relative 234expression changes of PpAHP5 after cold treatment (-28°C) in resistant and sensitive cultivars. (F) 235 Association between genotypes and cold hardiness (lethal temperature of 50%, LT50). (G) Allele 236 frequencies of association locus (Pp06: 9,187,362) in *PpAHP5* across seven groups.

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238 Next, we identified known biological processes that were overrepresented among 239 associations for each EV and for overall EVs (Supplementary Table S9). Functional categories 240 related to response to a series of abiotic or biotic stimuli, "programmed cell death (PCD)", "innate 241 immune response", and "LRR domain" were highly overrepresented (P < 0.05), suggesting that 242 EVs mainly shaped genomic regions related to stress responses. Notably, a series of processes 243 involved in secondary metabolism, including "flavonoid metabolic process", "jasmonic acid (JA) 244biosynthetic process", and "biosynthesis of plant hormones and terpenoids", were significantly 245 overrepresented (P < 0.05) (Supplementary Table S9). We found that genes related to JA 246 biosynthesis were enriched in altitude associations (P < 0.05). Previous studies have shown that 247 JA treatment contributes to enhanced cold resistance by promoting expression of the ICE-248 CBF/DREB1 transcriptional pathway, while a mutation in a key JA biosynthesis gene, LOX1 249 (Prupe.6G324400, an altitude association gene in this study), leads to cold hypersensitive 250 phenotypes (Hu et al. 2013). For each EV, several known biological processes were 251 overrepresented (Supplementary Table S9). For instance, genes involved in ion transport were 252highly enriched in those associated with soil pH (P < 0.05), as soil pH affects absorption of metal 253 ions in plants (Harter 1983).

254Temperature and precipitation are two of the most important EVs that affect plant 255distribution and survival. We identified temperature associated SNPs, distributed across all eight 256 peach chromosomes, and five association hotspots on chromosome 1, 2, 5, 6, and 8 were 257 detected in GWEAS for more than nine temperature-related EVs and altitude (Supplementary Fig. 258 S5A and 5B). Tolerance to low temperature in winter is a major factor that restricts the spread of 259 peach to extremely cold regions (north of 40 °N). To characterize genetic loci underlying 260 adaptation to extremely cold climates in peach, we performed a GWAS analysis of cold hardiness 261 and identified four association peaks, on chromosomes 2, 4, 6, and 7 (Fig. 2B). Of these, the peak 262 on chromosome 6 showed a strong selection signal, with sharp ROD in the NE group that 263 experienced an extreme cold winter (lowest temperature < -30 °C) (Fig. 2B). Moreover, this peak 264 overlapped with the temperature association hotspot on chromosome 6 and association peaks of 265 annual lowest temperature (Fig. 2B). The NE group (n = 19) inhabits areas north of 40 °N that 266 have extremely low winter temperatures, while the ST group (n = 14) grows in a contrasting 267 climate, south of 25 °N in areas with a warm winter (lowest temperature > 10 $^{\circ}$ C). We searched for 268 genomic regions and SNPs with extremely high differentiation between ST and NE groups. One 269 (Pp06: 9,187,362) of these SNPs (F_{ST} = 1) resided within the overlapping intervals between 270 annual lowest temperature and cold hardiness associations (Fig. 2C). This SNP was located in 271the gene PpAHP5 (Prupe.6G123100), belonging to a gene cluster encoding six histidine

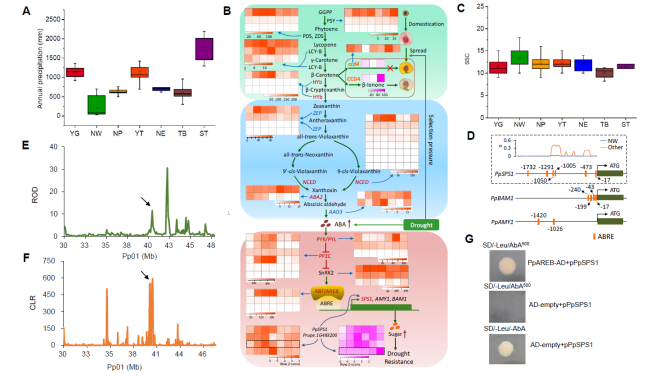
272 phosphotransfer proteins (AHP) (Fig. 2D), which have been reported to be involved in mediating 273 cold signaling in A. thaliana (Jeon and Kim 2013). Using cold treatment, we found this gene was 274 up-regulated by cold and resistance cultivars harbored significantly high expression level than 275sensitive one (Fig. 2E). At this SNP locus, all representative accessions in the NE group showed 276 a distinct genotype (TT) compared with the ST group (CC) (Fig. 2F), indicating that the TT 277 genotype in *PpAHP5* is favored in high-altitude cold regions (Fig. 2G), and that *PpAHP5* is a 278 candidate for conferring cold resistance in peach. We also detected six strong association regions 279 for precipitation-related EVs, including annual and seasonal precipitation, length of growing 280 season, aridity, and relative air humidity (Supplementary Fig. S5C and 5D). An extremely strong 281 association hotspot on Pp02 (4.8~7.2 Mb) was identified, exhibiting enrichments of R genes 282 (Verde et al. 2013), RLKs super family genes, NB-ARC domains, and other stress response-283 related genes, suggesting a genetic basis for precipitation adaptation.

284 To further elucidate the pattern of adaptation, we detected overlaps between selective 285 sweeps and GWEAS. A total of 888 genes (~23.3% of GWEAS genes) were shared between 286 selective sweeps and GWEAS (Supplementary Fig. S6). This revealed that although selective 287 sweeps are important, adaptations from standing variation or polygenic adaptation are also likely 288 an important mode of adaptation in peach, which may be related to its shortly spread history after 289 domestication (Li et al. 2019). These findings suggest that domesticated fruit species, such as 290 peach, are generating and enhancing adaptation by standing selection on existing multiple sites. 291 This situation is different from A. thaliana, which may have reached its adaptive limits owing to 292 the constraints imposed by the limited generation of new mutations (Hancock et al. 2011). 293 Collectively, these results indicate that both selective sweeps and GWEAS are central factors in 294 the adaptive genetics of domesticated species.

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Adaptation to highly drought regions

297 The NW group is from northwestern China, which has an extreme climate, characterized by 298 severe aridity (< 150 mm annual rainfall) (Fig. 3A) and extreme high or low temperatures in the 299 summer (> 40 °C) or winter (< -30 °C) (Supplementary Table S3). Peach accessions from this 300 region are frequently challenged by high drought stress. We found that genes overrepresented in 301 this group included those involved in abscisic acid (ABA) biosynthesis and signal transduction (P 302 < 0.05) (Supplementary Table S5), which are well known to regulate drought stress responses. 303 Transcriptome analyses of peach accessions grown under drought stress conditions revealed that 304 genes involved in the ABA pathway were highly enriched among differentially expressed genes 305 (DEGs), including NCED, PYR, ABA2, PP2C, and ABRE genes that showed selective signals in



306 the NW group (Fig. 3B), further suggesting a key role of ABA pathway in peach drought responses.

308 Fig. 3 Genetic basis of drought resistance and high sugar content in the NW group. (A) Annual 309 precipitation among the seven groups. (B) Relationship between the ABA pathway, drought stress and 310 evolution of flesh color. Heat map in orange indicate gene expression levels (FPKM) under drought 311 stress (0h, 6h, 12h, 24h, 3d, 6d, 12d). Heat maps in pink indicate gene expression levels (FPKM) 312 during peach fruit development (10, 50, and 90 days post bloom date (dpb) for PpCCD4; 20, 40, 60, 313 80, 100, 120 dpb for *PpSPS1*). Genes under selection in the NW group are highlighted in red. Red 314 arrows indicate the increase in levels of ABA and sugars. (C) Soluble solid content (SSC) among the 315 seven groups. (D) ABRE *cis*-acting elements in the promoters of *PpSPS1*, *PpBAM1*, and *PpAMY1*. 316 Orange boxes indicate ABRE elements in the promoter of each gene. The number around each ABRE 317 represents the position from the ATG. The distribution of ABRE elements and nucleotide diversity in 318 the promoter of *PpSPS1* in the NW and other groups are shown in a dashed box. (E) Distribution of 319 ROD around *PpSPS1* on chromosome 1. Black arrow points to *PpSPS1*. (F) Distribution of CLR values 320 around PpSPS1 on chromosome 1. Black arrow points to PpSPS1. (G) Verification of the interaction 321 between PpAREB (Prupe.1G434500) and the promoter of PpSPS1 (Prupe.1G483200) using a yeast 322 one-hybrid assay.

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324 Sugars function as the important signaling molecules in response to a range of abiotic and 325 biotic stresses in plants (Lastdrager et al. 2014). We found that peach fruits produced by 326 accessions from the NW group, especially accessions from Xinjiang province (Wang et al. 2012), 327 consistently had higher soluble sugar contents than those from other groups (Fig. 3C). Associated 328 long-term natural selection pressures contributing to greater accumulation of soluble sugars likely 329 include aridity, high diurnal temperature variation, and long sunshine duration. Moreover, the 330 starch and sucrose metabolism pathways were overrepresented in both DEGs under drought 331 stress treatment (35 genes) and genes under selection in the NW group (12 genes) (P < 0.05). 332 congruent with roles of sugars in drought stress. Furthermore, all the 12 genes in the selective 333 sweeps were differentially expressed following the drought stress treatment. We conclude that 334 higher soluble sugar contents in accessions from northwestern China represent an adaptive trait 335 driven by the local drought environment.

336 Previous studies of apple have demonstrated that drought stress and ABA contributed to 337 soluble sugar accumulation through the activation of sugar transporter and amylase genes by the 338 ABA-responsive transcription factor, AREB2 (Ma et al. 2017). Similarly, both drought stress and 339 exogenous ABA induce an increase in soluble sugar accumulation in peach fruit (Kobashi et al. 340 2000; Kobashi et al. 2001). Here we found that two putative gene targets of AREB2 (Fig. 3B and 341 3D), PpAMY1 (Prupe.1G142400) and PpBAM1 (Prupe.1G053800), were up-regulated by drought 342 treatment; however, neither exhibited a significant selection signal. To identify additional target 343 genes in drought mediated sugar accumulation, we searched for genes harboring the putative 344 binding domain of AREB2 among genes under selection in the NW group. This revealed a sucrose 345 phosphate synthase gene (PpSPS1, Prupe.1G483200), with six ABA-responsive elements 346 (ABREs) in the promoter region (Fig. 3D), showing a strong selection signal, with high ROD and 347 CLR values (Fig. 3E and 3F). *PpSPS1*, which is involved in the biosynthesis of sucrose, the 348 predominant soluble sugar in mature peach fruit and the key factor conferring sweetness, was 349 up-regulated by drought treatment (Fig. 3B), suggesting its roles in drought stress response. The 350 expression of *PpSPS1* increased by ~500-fold during fruit maturity (Fig. 3B), implying its roles in 351 fruit ripening and sugar accumulation. Using a yeast one-hybrid experiment, we verified the 352 interactions between AREB/ABF and the promoter of *PpSPS1* (Fig. 3G), providing new insight 353 into ABA-mediated enhanced sugar accumulation under drought stress. The selection on sugar 354 related genes may mediate adaptation to drought stress in the NW group, accompanied by the 355 increases in fruit sugar content. In addition, we found that the top of chromosome 5 and the middle 356 of chromosome 4, which have been reported to harbor major SSC- and sugar content-associated 357 guantitative trait loci (QTLs) and SSC candidate gene PpNCED3 (Martínez-García et al. 2013; Li 358 et al. 2019), also showed strong selection signals in the NW group. Selections on these genes 359 may underlie the genetic basis of high sugar levels in peach accessions grown in areas with high

360 drought stress. Moreover, such genes represent excellent candidates for high-sugar breeding.

361 Intriguingly, we found that flesh color of peach showed strong geographic pattern, with 362 ~80% of vellow-fleshed peach landraces originating from northwestern China (NW group). Yellow 363 flesh of peach mainly depends on the content of carotenoids at maturity, including β -cryptoxanthin 364 and β-carotene, and carotenoids are believed to be the major precursors for ABA biosynthesis 365 (Fig. 3B). A previous study has identified three loss-of-function variants involved in a carotenoid 366 cleavage dioxygenase gene (PpCCD4, Prupe 1G255500), leading to the abnormal carotenoid 367 degradation and yellow flesh (Falchi et al. 2013). The disturbed function of PpCCD4 in yellow-368 fleshed peach resulted in the retention of carotenoids, which can provide more precursors for ABA 369 biosynthesis (Fig. 3B), and may contribute to higher ABA levels and subsequent enhanced 370 drought tolerance. Moreover, using transcriptional analyses, we found that *PpCCD4* was down-371 regulated by drought treatments (Fig. 3B), suggesting its response to drought stress. Furthermore, 372 the carotenoid biosynthetic pathway was highly overrepresented with genes under selection in 373 the NW group (P < 0.05). Therefore, we conclude that yellow peach flesh has undergone long-374 term adaptive selection, driven by drought stress, and that modern yellow-fleshed peach cultivars 375 may originate from northwestern China.

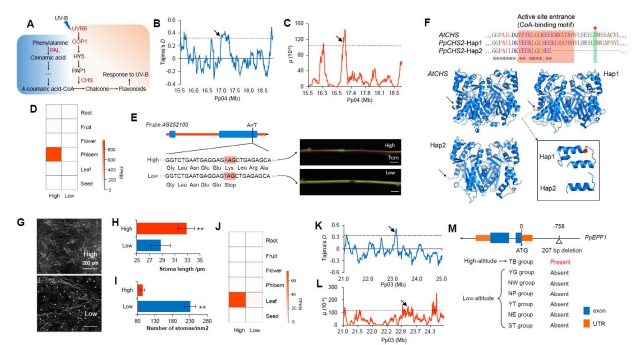
Collectively, we constructed a joint pathway for drought adaptation evolution in peach, driven by the complicated interactions between carotenoids, ABA, and sugar, of which ABA may be the central controller and play the key roles.

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380 Adaptation to high altitudes

381 Members of the TB group (n=45) are from 'the roof of the world', Tibet plateau, which is the highest 382 plateau on the earth, with an average elevation of 4500 m. This area is inhospitable to many 383 organisms because of its strong ultraviolet radiation, hypoxia, and severe cold (Supplementary 384 Table S3). At high altitudes, genome integrity is continuously challenged by intensive solar 385 ultraviolet radiation (UV-B, 280-315 nm)-induced DNA damage. Peach accessions in the TB group 386 tolerate these conditions using several adaptation-related phenotypes, such as a dark branch 387 color, epigeal germination, and red-colored new shoots (Supplementary Fig. S7). We identified 388 339 genomic regions, harboring 920 genes, showing signals of natural selection in the TB group 389 (Supplementary Table S4). Of which, we found a significant enrichment of genes involved in 390 'response to UV-B' category (P = 0.0004) (Supplementary Table S5), which is consistent with the 391 adaptation to high-altitude origin of the TB group. Flavonoids are a group of plant secondary 392 metabolites, which play important roles in UV-B protection (Li et al. 1993), and we found two 393 genes in the flavonoid biosynthetic pathway in the 'response to UV-B' category (Fig. 4A): chalcone

394 synthase (PpCHS2. Prupe.4G252100) and phenylalanine ammonia-lvase (PpPAL. 395 *Prupe.6G235400*), both of which showed strong selection signals in the TB group, with high µ and 396 Tajima's D values (Fig. 4B and 4C). Chalcone synthase catalyzes the first committed step in 397 flavonoid biosynthesis and previous studies showed that functional perturbation of an A. thaliana 398 homolog, AtCHS, resulted in UV-hypersensitive phenotypes, while in a UV-B-tolerant mutant 399 Atchs was up-regulated (Birza et al. 2001). We found that PpCHS2 was highly and specifically 400 expressed in the phloem of new shoots in the TB group (Fig. 4D), consistent with the red new 401 shoot phenotype. By scanning genomic variants in or around PpCHS2, we found that a SNP 402 (Pp04: 16,896,126, A>T) causing the introduction of a premature termination codon (Fig. 4E) 403 showed a high frequency in low altitude accessions (76.3%), but extreme low frequency of 404 substitution allele in the TB group (2.0%). Intriguingly, this SNP was located at the key active 405 region for protein function, CoA-binding motif (Fig. 4F), leading to an incomplete binding motif that 406 may result in the loss of function. Moreover, the premature termination resulted in the loss of one 407 conserved catalytic residue which is also crucial for catalytic activity (Ferrer et al. 1999). Therefore, 408 this SNP was designated as a candidate causative variant for the red new shoot phenotype 409 involved in flavonoid-mediated UV-B adaptation. Collectively, our results suggest that selection 410 on CHS gene and the regulation of anthocyanins may be one of important mechanisms to confer 411 avoiding damage from UV irradiation for peach at high altitudes.



412

Fig. 4 Genomic regions and candidate genes related to high-altitude adaptation in Tibet. (A)
 Pathway related to plant response to UV-B. Genes under selection are highlighted in red. (B-C)

415 Distribution of Tajima's D(B) and μ values (C) in the region around PpCHS2 (Prupe.4G252100) on 416 chromosome 4 (15.5-19.0 Mb). The dashed horizontal lines indicate a threshold of top 5% for Tajima's 417 $D \ge 0.36$) and μ test (≥ 1.07). Arrows point to *PpCHS2*. (**D**) Heatmap of expression profiles of *PpCHS2* 418 in different tissues in low- and high-altitude accessions. (E) A candidate stop-gained SNP in PpCHS2 419 that is associated with high altitude adaption and new shoot colors in accessions from low- and high-420 altitudes. (F) Effects of stop-gained SNP on protein structure of CHS. 3D structure of CHS protein was 421 obtained from Swiss-prot. The red shadow represents the CoA-binding motif. The green shadow 422 represents one of the conserved enzyme active site. (G) Scanning electron microscopy (SEM) of 423 stomata from the leaves of high- and low-altitude accessions. The magnification is 800x. (H-I) 424 Stomatal length (H) and stomatal density (i) in high- and low-altitude accessions. ** indicates P < 0.01. 425 (J) Heatmap of expression profiles of *PpEPF1* in different tissues in accessions from low- and high-426 altitudes. (K-L) Distribution of Tajima's D (K) and μ values (L) in a region around PpEPF1 427 (Prupe.3G235800) on chromosome 3 (21.0-25.0 Mb). The dashed horizontal lines indicate a threshold 428 of top 5% for Tajima's $D \ge 0.36$ and μ test ≥ 1.07). Arrows point to PpEPF1. (M) Structure of PpEPF1 429 and the position of the 207-bp deletion. The presence and absence of the 207-bp deletion in the seven 430 groups are given.

431

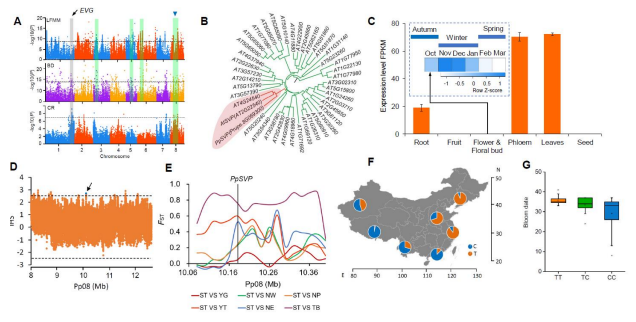
432 We observed that, compared with low-altitude accessions, those from high-altitudes had 433 a lower density of stomata and larger stomata size (Fig. 4G-4I). This may represent an adaptive 434 evolution to hypoxia at high altitudes. Interestingly, we found that the biological category 'stomatal 435 complex patterning' was significantly enriched in the gene set under selection (P = 0.008). By 436 transcriptional analyses of these genes, we found one of them, Prupe.3G235800, was highly and 437 specifically expressed in leaves, showing an altitudinal pattern with higher expression levels in 438 the TB group than in the low-altitude group (Fig. 4J). Notably, Prupe.3G235800, which encodes 439 the epidermal patterning factor 1 (PpEPF1) involved in stomatal development (Hara et al. 2009), 440 showed strong selection signals, based on the high Tajima's D and µ values (Fig. 4K and 4L). 441 Previous studies have shown that the mutation of a homolog of *PpEPF1* in *A. thaliana* results in 442 increased stomatal density (Hara et al. 2009). By scanning the variants in *PpEPF1*, we found that 443 SNPs with functional significance were absent. Through further scanning variants at the upstream 444 or downstream of *PpEPF1*, we identified a TB group specific 207-bp deletion in the promoter 445 region (-758 bp from the start codon) of *PpEPF1* (Fig. 4M), suggesting that the adaptive evolution 446 controlled by *PpEPF1* may be mediated by regulation of its expression. Furthermore, over-447 expression of *PpEPF1* in *A. thaliana* resulted in a decrease in stomatal density (Supplementary 448 Fig. S8). These findings suggest that selection on *PpEPF1* may be closely related to adaptation 449 to hypoxia in high-altitudes through the regulation of stomatal density.

450

461

451 A major SVP locus involved in adaptive evolution of bloom date

452 Bloom date (BD) is crucial for local adaptation in peach, and is controlled by multiple genes (Fan 453 et al. 2010). To explore the genetic basis of adaptation of BD, we performed GWAS of BD using 454 174 accessions that were phenotyped. This revealed 399 associated SNPs and 12 association 455 peaks (Fig. 5A), of which six overlapped with previously reported QTLs (Fan et al. 2010). Next, 456 we identified candidates involved in local adaptation by detecting SNPs showing associations with 457 EVs using a latent factor mixed-effect model (LFMM), resulting in a final set of 23 association 458 peaks (Fig. 5A). By overlapping BD GWAS and LFMM analyses, we found four regions on 459 chromosomes 3, 5, 6, and 8 that may underlie the local adaptation of BD during spread of peach 460 to different climates (Fig. 5A).



462 Fig. 5 A major PpSVP locus involved in local adaptation of bloom date in peach. (A) Manhattan 463 plots of SNPs associated with EVs (LFMM), bloom date (BD), and chilling requirement (CR). Dashed 464 lines represent the significance thresholds for the tests. The overlapped regions between GWAS for 465 BD and LFMM are highlighted using green shaded rectangles. The major QTL for CR and BD 466 overlapping with local selection signals on chromosome 8 surrounding *PpSVP* is indicated by a blue 467 triangle. The EVG locus is highlighted using a gray shaded rectangle. (B) Neighbor-joining tree of 468 *PpSVP* and MIKC-type MADS family genes. The clade containing *PpSVP* is highlighted in red. (C) 469 Temporal and spatial expression patterns of *PpSVP*. Error bars represent standard deviation of three 470 biological replicates. (D) Patterns of normalized iHS scores across the ~4 Mb genomic region around 471 PpSVP. The dashed horizontal lines represent the threshold of positive selection signal (|iHS| > 2.5). 472 The blue dot indicates the SNP (Pp08: 10,173,576) that showed high iHS score in PpSVP. (E) F_{ST} 473 around *PpSVP* among different groups. The associated SNP in *PpSVP* is indicated using vertical black

- Ine. (F) Allelic frequencies of the associated SNP (Pp08: 10,173,576) in *PpSVP* across seven groups.
- 475 (G) Relationship between genotypes of associated SNP (Pp08: 10,173,576) and bloom date.
- 476

477 Chilling requirement (CR) is another important adaptive trait and is significantly correlated 478 with BD. We re-performed the GWAS for CR based on our previous study (Li et al. 2019) using 479 174 landrace accessions and identified six association peaks, of which three (chromosome 1, 7, 480 and 8) were shared with BD (Fig. 5A), including the major QTL for CR harboring the EVG locus 481 conferring dormancy mutation in peach (Li et al. 2009). After overlapping GWAS of CR and BD 482 with the LFMM analysis, we found a strong overlap spanning ~4-Mb on chromosome 8, which 483 may be important for local adaptation of BD in peach (Fig. 5A). Interestingly, the major QTL for 484 CR and BD on chromosome 1 showed no local adaptation signal in the LFMM analysis (Fig. 5A), 485 suggesting that climates may drive the evolution of BD and CR by shaping QTLs with small effects.

486 The 4-Mb region encompasses 275 genes, including a putative ortholog of A. thaliana 487 SHORT VEGETATIVE PHASE (PpSVP, Prupe.8G069300). SVP is involved in controlling 488 flowering time and has previously been implicated in regulating dormancy in *Prunus* (Li et al. 2009; 489 Sasaki et al. 2011; Zhang et al. 2012). Phylogenetic analysis confirmed that *PpSVP* belongs to a 490 MADS-box family and is closely related to the AGL22 subfamily (Fig. 5B). PpSVP showed strong 491 tissue-specific expression, with high expression only in vegetative organs. Moreover, expression 492 of *PpSVP* was up-regulated during dormancy induction and down-regulated by winter chill (0-493 7.2 °C) and by forcing temperature (heat) in floral buds in spring (Fig. 5C), suggesting its potential 494 role in regulating BD and CR. Moreover, through calculating the standardized integrated 495 haplotype score (iHS) for SNPs located in this overlap region, we found a strong positive selection 496 signal around the *PpSVP* locus (Fig. 5D). Additionally, an exceptionally high F_{ST} value was 497 identified in this region, especially between the ST and NE groups and between the ST and YT 498 groups (Fig. 5E) that harbor distinct bloom date. The *PpSVP* locus thus represents a strong 499 candidate gene for local adaptation of BD and CR. We propose that spatially varying selection 500 has driven latitudinal differentiation at this locus. Positive selection signals, revealed by a CLR 501 test, were also detected in the NE and ST groups (Fig. 5F). Overall, all these results provide 502 compelling evidence of local selection on the PpSVP locus during adaptive evolution to different 503 climates after domestication.

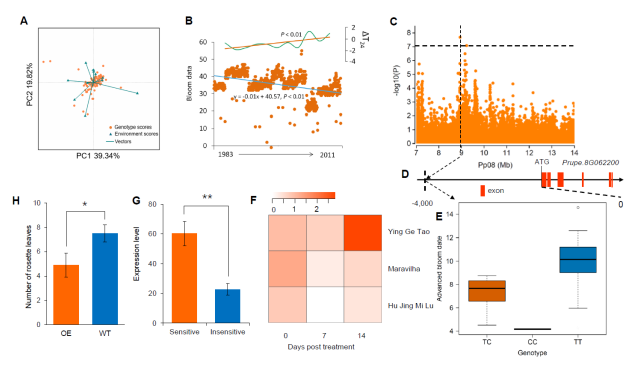
504 To identify the causal variants underlying adaptation of BD, we screened for SNPs with 505 high F_{ST} between the NE (late bloom) and ST (early bloom) groups at the *PpSVP* locus. No SNP 506 with high differentiation was identified that caused an amino acid change. However, a SNP located 507 at 5'-untranslated regions (5'-UTR) with high F_{ST} value (F_{ST} =0.9) was identified, suggesting that 508 the BD and CR may adapt to different climates through shaping the expression of the controlled 509 gene. Allele frequencies of this SNP showed strong geographical pattern and the early bloom 510 alleles (CC) mainly occurred in low altitude regions (ST and YG groups) and wild group (TB group) 511 (Fig. 5G and 5H), consistent with phenotype. This also provides insights into two distinct 512 evolutionary routes of BD and CR to low and high chill regions. Moreover, overexpression of the 513low-altitude favored genotype of PpSVP (CC) in A. thaliana resulted in plants with strong 514 vegetative growth and delayed flowering time (Li et al. 2019).

515

516 Genomic locus associated with response to climate change

517 Adaptation to accelerating rates of climate change is increasingly important for species survival. 518 The advance in bloom date (ABD), as a consequence of global warming over recent decades, 519 has been observed in many temperate species, including peach (Menzel et al. 2006; Li et al. 520 2016). However, the genetic mechanism underlying ABD have not been characterized. We 521 performed a long-term observation of BD with 89 peach accessions spanning three decades, from 522 the 1980s to 2010s (Supplementary Fig. S9A). We observed a significant ABD (P < 0.001), based 523 on an additive main effects and multiplicative interaction (AMMI) analysis (Annicchiarico, 1997). 524 and the main driver was found to be a temperature rise in the spring (explained 61.3% variation, 525 P < 0.001) (Fig. 6A). Using a linear regression analysis, we quantified ABD and found that BD 526 has advanced by approximately 10 days on average over last 30 years (Fig. 6B).

527 Next, we performed GWAS for ABD to identify genetic loci associated with responses to 528 global warming (Supplementary Fig. S9B). This revealed a strong association peak on 529 chromosome 8 ($P < 7.28 \times 10^{-8}$) (Fig. 6C) in an area harboring 14 candidate genes around peak 530 association. This association was also located at overlap among GWAS signals of CR, BD and 531LFMM analysis. The most significant SNP was located in a region upstream of *Prupe*.8G062200, 532 with genotype of TT showing sensitive to global warming and CC insensitive (Fig. 6D and 6E). 533 *Prupe.8G062200* encodes a putative night light-inducible and clock-regulated 1 (LNK1) protein, 534and showed high expression levels at blooming. A homolog of this gene in A. thaliana is involved 535 in regulation of the circadian clock, which regulates COL1 genes at warm temperatures, and thus 536 a potential regulator of flowering time (Mikkelsen and Thomashow 2009; Rugnone et al. 2013). A 537 simulation experiment showed that *PpLNK1* was up-regulated by rising temperatures during heat 538accumulation, suggesting that *PpLNK1* may be up-regulated by temperature rise in spring (Fig. 539 6F). In addition, expression of *PpLNK1* in peach accessions that are sensitive to global warming 540 was significantly higher than in those that are insensitive (Fig. 6G). Notably, over-expression of 541 PpLNK1 in model plant, A. thaliana, led to the early flowering (Fig. 6H). Moreover, several cis-542 elements associated with temperature and light responsiveness was identified (Supplementary 543 Table S10). Therefore, we conclude that *PpLNK1* may play important roles in regulating annual 544 circadian clock of flowering time as influenced by rising temperature in peach. PpLNK1 is thus a 545 plausible candidate gene for responses to global warming, but further work will be necessary to 546 provide more direct evidence of its roles. Collectively, our comprehensive analyses detected 547 genomic loci associated with responses to global warming, which can improve our understanding 548 of the genetic architecture of plant adaptation to global climate change.



550 Fig. 6 Genotype-environment interaction analysis and genome-wide association study of 551advance in bloom date. (A) Genotype-environment interaction analysis of bloom date from 1983 to 552 2011 using the AMMI analysis. (B) Scatter plots of relative bloom date of 89 peach accessions from 553 1983 to 2011 and temperature change in the spring. The blue and orange lines represent the trend of 554bloom date changes and temperature changes in the spring, respectively, based on the linear 555 regression analyses. ΔT24 indicates anomalies in the mean temperature from February to April 556 compared to those from 1983-2011. (C) Regional Manhattan plot of GWAS for ABD on chromosome 557 8 of the 7.0-14.0 Mb region. The gray dashed line indicates significance threshold ($P < 7.28 \times 10^{-8}$ or -558 log10(P) > 7.08) using a Bonferroni test (0.05). (**D**) Most significant SNP associated with ABD and its 559 location relative to gene PpLNK1 (Prupe.8G062200). (E) Association between genotypes of the most 560 significant SNP and ABD. (F) Changes in *PpLNK1* expression in three cultivars in a climate warming 561 simulation experiment. dpt, days post treatment. (G) Comparison of PpLNK1 expression between

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accessions sensitive and insensitive to global warming at blooming. ** represents P < 0.01. (H) 563 Comparison of BD between wild type (WT) and *PpLNK1* over-expression (OE) *A. thaliana* lines. * 564 indicates P < 0.05.

Long-term observation of BD enabled multi-year GWAS. We identified a total of 713 SNPs associated with BD ($P < 7.28 \times 10^{-8}$), including 483 temporary associations that were identified only in one year, 214 associations in at least two years, and 16 stable associations in more than five years, of which several overlapped with previous reported QTLs (Fan et al. 2010) (Supplementary Table S11). Among stable associations, a strong association peak within a small intergenic region (Pp06: 15,327,714~15,354,080) on chromosome 6 was identified in eight years of GWAS, which can be further developed for marker-assisted selection.

572

573 Conclusions

574 Plant genomes have been shaped by natural selection during the local adaptation to diverse 575 environmental conditions. Peach provides an excellent model to investigate the genetic basis and 576 mode of adaptation to climate change, thanks to its relatively small genome size (~227.4Mb) and 577 extensive climatic variation across its native range. We generated a large variation map for peach 578 through sequencing of a climate-extensive panel of 263 peach landraces and wild relatives. 579 Notably, we first detected the genetic basis of adaptation to high altitudes for fruit species, P. mira 580 (TB group), and we found that genes involved in the biosynthesis of flavonoids (*PpCHS2*) and 581 stomatal development (*PpEPF1*) may play important roles in overcoming strong UV-B radiation 582 and hypoxia, respectively, on the Tibet Plateau. We discovered that high sugar content and yellow 583 flesh of peach in drought regions were drought-induced adaptive evolution mediated by 584 interactions between the abscisic acid pathway, *PpSPS1* and carotenoids. More than nine 585 thousand genomic loci, associated with 51 specific climate variables, were identified. These 586 included several hotspots associated with temperature and precipitation, as well as a SNP 587 associated with cold hardiness. Integrative analyses of selective sweeps and GWEAS suggest 588 that peach adaptation was generated and enhanced by standing selection on multi sites. Genomic 589 loci underlying the local adaption of BD and CR were found to be two evolutionary adaptations to 590 low and high latitude regions. In addition, through data collected over a 30-year period, we 591 identified a candidate genetic locus associated with responses to global warming in plant species.

592 This study provides new insights into peach adaptation to its habits and how climate has 593 shaped the genome of a perennial tree plant through natural selection. These results also provide 594 a new resource for studies of peach evolutionary biology and breeding, especially with regard to 595 enhancing stress-resistance.

596 Methods

597 Plant materials and sequencing

598 A total of 263 peach accessions were sampled from the NPGRC (National Peach Germplasm 599 Repository of China), except the 45 P. mira accessions, which were sampled from the Tibet 600 plateau. These accessions, collected from almost all the distribution regions of peach landraces 601 and wild relatives, including seven major ecotypes. These accessions included 45 of P. mira 602 Koehne, 4 of P. davidiana (Carr.) Franch., 2 of P. kansuensis Rehd., a single P. potaninii Batal., 603 205 of P. persica L., and 6 of P. ferganensis Kost. et Riab (Supplemental Table 1). Of these, P. 604 persica L. and P. ferganensis Kost. et Riab accessions belong to landraces, while the others are 605 wild relatives. Total genomic DNA was extracted from young leaves using the 606 cetyltriethylammnonium bromide (CTAB) method (Murray and Thompson 1980). At least 4 µg of 607 genomic DNA from each accession was used to construct pair-end sequencing libraries with 608 insert sizes of approximately 300-bp or 500-bp following the manufacturer's instructions (Illumina 609 Inc.) (Supplemental Table 1). A total of >1 Gb of sequence data was generated for each accession 610 from 49-bp, 90-bp, or 125-bp paired-end reads, using the Illumina GA or HiSeg 2500 platform 611 (Illumina, San Diego, USA) (Supplemental Table S1).

612

613 Read mapping and variation calling

614 Pair-end reads from each accession were mapped to the peach Lovell genome (release v2.0) 615 using BWA (Li and Durbin 2009) (Version: 0.7.12) with the following parameters: bwa mem -t 4 -616 M -R. Read alignments were converted into the BAM format, sorted according mapping 617 and PCR duplicates Picard coordinates. removed using the package 618 (http://broadinstitute.github.io/picard/; Version: 1.136) with default parameters. The coverage and 619 depth of sequence alignments were computed using the Genome Analysis Toolkit (GATK, version: 620 3.4-46; see URLs) DepthOfCoverage program (McKenna et al. 2010). The coverage and depth 621 of each accession are detailed in Supplemental Table S1.

To accurately identify SNPs, the low-quality alignments (a mapping quality score <20) were filtered using SAMtools (Li et al. 2009). SNP detection was performed using GATK HaplotypeCaller, which identifies SNPs by local *de novo* assembly of haplotypes in an active region (Depristo et al. 2011). The detailed processes were as follows: (1) After filtering the lowquality alignments, the reads around the INDELs were realigned through two steps, including identifying regions where realignment was needed using the GATK RealignerTargetCreator package, and realigning the regions found in the first step GATK IndelRealigner package. Next, 629 a realigned BAM file for each accession, which was used for SNP detection, was generated using 630 GATK PrintReads packages. (2) SNPs were detected at a population level using the realigned 631 BAM file with GATK HaplotypeCaller. To reduce the number of false positives, a high SNP 632 confidence score was set with the following parameters: -stand_call_conf 30 -stand_emit_conf 633 40. (3) To ensure the quality of variant calling, a hard filter was applied for the raw SNPs with 634 SNP quality > 40 and the number of supporting reads > 2, using GTAK VariantFiltration, with the 635 following parameters: QUAL < 40, QD < 2.0, FS > 60.0, MQ < 40.0, MQRankSum < -12.5, 636 ReadPosRankSum < -8.0, -cluster 3, -window 10.

637 The accuracy of SNPs was assessed using a Sequenom MassARRAY platform (Sequenom, 638 San Diego, USA), following the manufacturer's protocol. A total of 18 randomly selected SNPs 639 was investigated in 130 accessions. The list of accessions is provided in Supplemental Table S2. 640 INDEL calling was performed using the same pipeline as the SNP calling since the GATK is 641 capable of calling SNPs and INDELs simultaneously. To reduce the number of false positives, we 642 also applied a harder filter for raw INDELs using GTAK VariantFiltration with the following 643 parameters: QD < 2.0, FS > 200.0, ReadPosRankSum < -20.0. Insertions and deletions ≤6 bp 644 were defined as the small INDELs.

645 SV calling was performed using the SpeedSeg (Chiang et al. 2015), DELLY (Tobias et al. 646 2012), and manta (Chen et al. 2016) programs. For SpeedSeg calling, paired-end reads were 647 mapped to the reference genome using the 'align' module in SpeedSeg and the following 648 parameters: speedseq align -R -t 4. Three BAM files were generated, including a full, duplicate-649 marked, sorted BAM, a BAM file containing split reads, and a BAM file containing discordant read-650 pairs. SVs were identified using the 'sv' module in SpeedSeg, using the following settings: 651 speedseg sv -o -x -t 25 -R -B -D -S -g -P. For DELLY calling, mapped pair-end reads in BAM 652 format, generated by BWA-MEM (Li and Durbin 2009) after sorting and marking PCR duplicates, 653 were used as input files. SVs were identified using the call module in DELLY with default 654 parameters. SV files in VCF format for all of 263 samples were merged into a population level 655 VCF file using BCFtools (Li et al. 2009). For SV calling with manta, the same BAM files with 656 DELLY were used to detect SVs, with default parameters. SV files for 263 accessions were then 657 merged using SURIVAR (Jeffares et al 2017) and genotyped using SVtyper (Chiang et al. 2015) 658 with default parameters. Finally, SVs identified by at least two callers were designated as the final 659 set of SVs.

660

661 SNP annotation

662 SNP annotation was performed based on genomic locations and predicted coding effects,

according to the peach genome annotation (release annotation v2.1, see URLs), using the snpEff (Cingolani et al. 2012) (Version: 4.1g). The final SNPs were categorized in exonic regions, intronic regions, splicing sites, 5' UTRs and 3' UTRs, upstream and downstream regions, and intergenic regions, based on the peach genome annotation. SNPs in coding sequence were further grouped into synonymous SNPs (no amino acid changes) and nonsynonymous SNPs (amino acid changes). SNP effects were further divided into four types according to their impacts on gene function, including HIGH, MODERATE, LOW, and MODIFIER.

670

671 **Population genetics analysis**

672 To build a phylogenetic tree, we selected a subset of 2,468,307 SNPs with minor allele frequency 673 (MAF) >0.05 in all 263 accessions from the final SNP data set (4,611,842). A neighbor-joining 674 tree was constructed using PHYLIP (Felsentein 1989) (Version: 3.696) on the basis of the distance 675 matrix with 1,000 The bootstrap replicates. software FigTree 676 (http://tree.bio.ed.ac.uk/software/figtree/; version: 1.4.2) was used to visualize the neighbor-677 joining tree. The principal component analysis (PCA) was performed based on the same SNPs 678 data set (2,468,307 SNPs with MAF > 0.05) using the smartpca program in the EIGENSFOT⁷⁰ 679 software (Version: 6.0.1) with default settings (Price et al. 2006). The first three eigenvectors were 680 used to plot the data in two and three dimensions. The population structure was also investigated 681 using the same SNP data set (2,468,307 SNPs with MAF>0.05) with the FRAPPE (Version: 1.1) 682 software (Tang et al. 2005), which is based on a maximum likelihood method. We ran 10,000 683 iterations, and the numbers of clusters (K) were set from 2 to 8.

684

685 Identification of select sweeps

686 To detect signals of selective sweeps, we selected three distinct genome-wide selection metrics 687 for each group (excluding the TB group), including the reduction of nucleotide diversity (π) , 688 Tajima's D, and genetic differentiation (F_{ST}). We calculated these three selection metrics based 689 on all SNPs (4,611,842) using VCFtools (Danecek et al. 2011) (Version: 0.1.13), with a 10-kb 690 window and a step size of 1 kb. We defined the empirical top 5% of windows or regions as 691 candidate selective outliers for each selection scan metric. The adjacent selective outliers were 692 merged. For each population, selection outliers detected in at least two of the selection scan 693 metrics were designated as the candidate selection regions (CSRs). The TB group consisted of 694 wild relatives (P. mira) and three other methods were used to detect selective sweeps: Tajima's 695 D, RAiSD (Alachiotis et al. 2018), and CLR (Pavlidis et al. 2013). Similarly, the top 5% of windows 696 or regions identified in at least two metrics were designated as candidate selective sweeps.

697 **Collection of climate variables**

698 A total of 51 environmental variables were selected as being essential for peach growth and 699 survival (Supplemental Table S6), representing extremes and seasonality of temperature and 700 precipitation, altitude, latitude, relative air humidity, water vapor pressure, growing season lengths, 701 and aridity. Of these, 39 datasets of climate variables were downloaded from WorldClim 702 (http://www.worldclim.org; version: 1.4), with a resolution of 2.5 minutes, and climate variables for 703 each accession were extracted using DIVA-GIS (http://www.diva-gis.org; version: 7.5) 704 (Supplemental Table S6). Six climate variables were downloaded from CDMC 705 (http://data.cma.cn/en/?r=site/index) and climate variables for each accession were extracted 706 using ArcGIS (http://www.arcgis.com; version: 10.3) (Supplemental Table S6). Four climate 707 variables were downloaded from the FAO (http://www.fao.org/geonetwork/srv/en/main.home), 708 with a resolution of 5 minutes or 10 minutes and climate variables for each accession were 709 extracted using ArcGIS (Supplemental Table S7). Altitude and latitude for each accession were 710 recorded using a GPS (Magellangps triton 300E; http://www.magellangps.com) when the 711 accessions were collected.

712

713 Genome-wide environmental association study (GWEAS)

714 GWEAS was performed for 51 climate variables using 4,611,842 high-guality SNPs. The 715 association analyses were performed using the mixed linear model (MLM) with Efficient Mixed-716 Model Association eXpedited (EMMAX) software (Zhou and Stephens 2012). To minimize the 717 number of false positives and increase statistical power, population structure was corrected using 718 a kinship matrix, which was estimated with EMMAX emmax-kin program (Zhou and Stephens 719 2012). The genome-wide significance thresholds of the GWEAS were determined using the 720 Bonferroni test. Based on a nominal level of 0.05, the threshold was set as 0.05/total SNPs 721 $(\log 10(P) = -7.13).$

722

723 **Functional enrichment and pathway analysis**

To test whether candidate genes were overrepresented among lists from known biological processes, gene families and pathways, a functional enrichment and pathway analysis was performed based on Fisher exact tests (P < 0.05), using the Database for Annotation, Visualization and Integrated Discovery (DAVID) (Huang et al. 2009) (Version: 6.7). To obtain the comprehensive functional annotations, a list of annotation categories was selected, including GO terms and KEGG pathway. The annotation analysis was performed for genes that were in selective sweeps and GWEAS associations.

731 Phenotyping and genome-wide association study (GWAS)

732 The first bloom date (BD) was measured at the National Peach Germplasm Repository of China 733 (NPGRC) (N34.71°, E113.70°, A.S.L. 74 m), located in Zhengzhou, Henan Province, China. The 734 first bloom date data used span February 25 to April 25 from 1983 to 2011 as this period captured 735 the majority of diversity of BD. A total of 89 accessions, with each represented by two replicates, 736 were used to investigate BD (Supplemental Fig. S9A). The first bloom date was defined as the 737 day when approximately 5% of the flowers have completely opened. The advance in bloom date 738 (ABD) for each accession was estimated using a linear regression analysis, based on the BD 739 from 1983 to 2011. The ABD information for each accession is detailed in Supplemental Fig. S9B.

740 To identify genetic loci associated with ABD, GWAS was performed for ABD based a set of 741 873,895 SNPs, identified after removing SNPs with low-frequency (MAF < 0.05) and a high 742 missing rate (missing rate > 0.2) using the EMMAX program (Zhou and Stephens 2012). To 743 minimize the number of false positives and to increase the statistical power, population structure 744 was corrected using a kinship matrix, which was calculated with EMMAX emmax-kin program 745 (Zhou and Stephens 2012). The genome-wide significance threshold of the GWAS was 746 determined using the Bonferroni test. Based on a nominal level of 0.05, the threshold was set as 747 0.05/total SNPs (log10(P) = -7.08). GWAS was also performed for yearly BD data from 1983 to 748 2011 based on the same SNP data set, using the same method as above.

749 For CR, phenotyping analyses were performed in 2011 and 2012 as in Fan et al (2010). A 0-750 7.2°C model was chosen to evaluate CR and the number of hours in this range (chilling hours; 751 CHs) was counted, starting when the daily average air temperature dropped to below 7.2°C. 752 Starting at 50 CHs, the branches of each accession were cut every 50 CHs until 1,300 CHs. For 753 each accession, two clones were sampled, and three branches longer than 40 cm with floral buds 754 were taken from each clone. Branch cuttings were placed in water in a greenhouse at 25°C and 755 a 16 h/8 h photoperiod to force floral bud break. The frequency of floral bud break was evaluated 756 after two weeks. The CR of an accession was defined as being sufficient at a specific sampling 757 time if 50% of floral buds on the branch cuttings opened. GWAS for CR was also performed using 758 MLM in EMMAX.

Cold hardiness was evaluated using a conductance-based semi-lethal temperature method in December-January of 2013-2014 on 143 accessions. Six annual branches longer than 20 cm were sampled for each accession. A total of six cold treatments were used: -10, -15, -20, -25, -30, and -35°C. Branch cuttings were incubated in freezer with the six treatments for 24 h. After cold treatments, the cuttings were placed at 0°C for 8 h. Branch cuttings were then cut into 2 mm segments. A total of 2 g of segments was used to measure the conductance, with three biological replicates. The initial conductance (C1) was measured after a 12 h steep in 10 ml water. The final

conductance (C2) was measured after boiling the samples for 20 min and leaving them to cool to

room temperature for a subsequent 2 h period. The relative conductance (RC) was calculated

vising following formula:

769 RC=(C1/C2) × 100

Finally, the semi-lethal temperature (LT50) was calculated using a logistic function based onRC.

772

773 Yeast one-hybrid assay

Yeast one-hybridization assay was performed using the Matchmaker[®] Gold Yeast One-Hybrid System (Clontech, Palo Alto, CA, USA). The promoter sequence (upstream 2kb from the start codon) of the sucrose phosphate synthase, *PpSPS1 (Prupe.1G483200*), was cloned into the pAbAi vector. Similarly, the full-length of ABA-responsive element binding 1, *PpAREB1* (*Prupe.1G434500*), was subcloned into the pGADT7 AD vector. The auto-activation and TF– protein interaction analyses were conducted according to manufacturer's protocol.

780

781 Scanning electron microscopy (SEM)

782 Stomata were examined by SEM in young leaves from three accessions from the TB group and 783 three accessions from the NP group, representing high-altitude and low-altitude accession, 784 respectively. Three replicates were sampled from each accession. Samples were fixed in 2.5% 785 glutaraldehyde (pH = 7.4) for 24 h at 4° C. Subsequently, fixed samples were dehydrated using an 786 ethanol series (30% ethanol, 20 min; 50% ethanol, 20 min; 70% ethanol, 20 min; 100% ethanol, 787 30 min (twice)). The dehydrated samples were then dried in a critical-point drving apparatus 788 (Quorum K850; England). Dried samples were mounted on stubs and sputter-coated with gold 789 (FEI; America) and observed under a scanning electron microscopy (SEM) (FEI Quanta 250; 790 America).

791

792 RNA-Seq analysis

For drought stress treatment, four-year-old potted peach seedlings from peach cultivar "Dong Xue Mi Tao" were used. Fruit flesh were taken at six drought stress treatment time points, including 6 hours, 12 hours, 24 hours, 3 days, 6 days, and 12 days. For expression profiles in different tissues, roots, leaves, fruit, flowers, phloem, and seeds were sampled from "Aba Guang He Tao" (highaltitude) and "B-4" (low-altitude). For the expression of *PpCCD4*, fruit fleshes were sampled from "Zao Huang Pan Tao" (yellow-fleshed) and "Zhong Tao Hong Yu" (white-fleshed) at 10, 50, and

799 90 days post bloom date (dpb). For the expression of *PpSPS1*, fruit fleshes were sampled from 800 "Chinese cling" at 20, 40, 60, 80, 100, 120 dpb. Three biological replicates were collected for each 801 sample. The tissues were immediately frozen in liquid nitrogen and then ground to fine powder. 802 Total RNA was extracted using a quick extraction kit (Aidlab, Beijing, China). First and second 803 strand complementary DNA (cDNA) was synthesized using a cDNA Synthesis System kit 804 (TOYOBO, Osaka, Japan), following the manufacturer's protocol. Double-strand cDNAs were 805 then purified and adapters were ligated to the short fragments. The constructed RNA-Seq libraries 806 were sequenced using the Illumina HiSeg 2000 platform (Illumina, San Diego, USA) in paired-807 end 150-bp mode. Low-quality reads were filtered from the raw reads using Trimmomatic (Bolger 808 et al. 2014). Data analysis followed the protocol proposed by Pertea et al (2016). Cleaned reads 809 were mapped to the peach reference genome using Hisat2 (Version 2.0.5) (Kim et al. 2015) with 810 default parameters. Transcript abundances were calculated and transcript assembly was 811 performed using Stringtie (Pertea et al. 2015). DEG analysis was carried out using the R package 812 ballgown (Frazee et al. 2015).

813

814 Over-expression of candidate genes in *A. thaliana*

815 The full-length open reading frames of three peach genes, *PpEPF1* (*Prupe.3G235800*), *PpSVP* 816 (Prupe.8G069300), and PpLNK1 (Prupe.8G062200), were amplified by PCR using cDNAs 817 derived from young leaves of "Aba Guang He Tao", 'Nanshan Tian Tao' (CR=200h), and 818 "Nanshan Tian Tao", respectively. The PCR products were cloned into the pBI121 vector driven 819 by the cauliflower mosaic virus (CaMV) 35S promoter at Sangon Biotech (Sangon, Shanghai, 820 China). The resulting constructs were then transformed into A. thaliana Columbia type using 821 Agrobacterium tumefaciens GV3101 and positive transformants selected with kanamycin. Ten 822 transgenic lines for each gene were used to evaluate the flowering time. The stomata size and 823 density were observed under a light microscope (Olympus BX51, Tokyo, Japan) with a 400 x 824 objective lens.

825

826 **RNA extraction and expression analysis using qRT-PCR**

For *PpSVP* expression analysis, floral buds from 'Nanshan Tian Tao' were sampled on October 15, November 15, December 15, January 15, February 15, March 15 in 2016-2017. *PpLNK1* expression was measured in floral buds (blooming soon) from three global warming-sensitive accessions ('Wu Yue Xian', 'Nanshan Tian Tao', and 'Li He Pan Tao') and three global warminginsensitive accessions ('Xinjiang Pan Tao', 'Wuhan 2', and 'Kashi 2') at 2016 and 2017. For *PpAHP5*, the phloem (including cambium) was collected from two cultivars 'Hunchun Tao' (cold 833 resistant) and 'Nanshan Tian Tao' (cold sensitive) after 24 hours treatment under -28 °C 834 refrigerator and following 21 °C incubation in water. For each sample, three biological replicates 835 were used. Total RNA was extracted using an extraction kit (Aidlab, Beijing, China) and first-836 strand cDNA was synthesized with 1µg RNA using a FastQuant RT Kit (with gDNase) (TIANGEN, 837 Beijing, China). Gene-specific primers were designed using Primer-BLAST software (National 838 Center for Biotechnology Information, Maryland, USA). qRT-PCR was performed using a SYBR 839 green I master kit (Roche Diagnostics, Indianapolis, USA) with the LightCycler System (Roche 840 LightCycler 480, Indianapolis, USA), following the manufacturer's protocol. Relative expression levels were calculated using the $2^{-\Delta\Delta CT}$ method. A β -actin was used as the reference gene. 841

842

843 **Global warming simulation experiment**

844 The global warming simulation experiment was performed in 2016-2017. Three peach cultivars 845 (Nanshan Tian Tao, Hu Jing Mi Lu, and Maravila), each with two clones, were used as plant 846 materials. For each cultivar, ~30 annual branches longer than 40 cm with floral buds were taken 847 from each clone when the winter chill accumulation was ~900 chilling hours (0~7.2°C, excluding 848 0°C). Branch cuttings were placed in water in greenhouse at 25°C and with a 16 h/8 h photoperiod. 849 to simulate climate warming. The ratio of bud break was investigated daily, starting from the day 850 that the branch cuttings were placed in the greenhouse. The floral buds, excluding the 851 tegmentum, were collected weekly and frozen in liquid nitrogen. The sampled floral buds were 852 used for qRT-PCR analyses following the protocol described above.

853

854 Data access

- 855 Raw sequence data have been deposited in the NCBI Short Read Archive (SRA) under
- accession SRP108113. SNPs and SVs in Variant Call Format (VCF) have been deposited into
- 857 the Figshare database (SNPs:
- 858 https://figshare.com/articles/SNPs_for_263_peach_accessions/7636715, SVs:
- 859 https://figshare.com/articles/SVs_for_peach_sequencing/7636721). All other relevant data are
- 860 contained within the paper and available in supplementary files.
- 861

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DNA samples; Y.L., K.C., and N.L. performed the data analyses; Y.L., T.D., J.W., L.G., Q.H., and
W.F. performed phenotyping. Y.L. performed the molecular experiment. Y.L. and K. C. wrote the
paper; L.W., Z.F., W.G., and S.H. revised the paper. All authors read and approved the final
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