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Genome-wide view of genetic diversity reveals paths of selection and cultivar differentiation in peach domestication

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Abstract
Domestication and cultivar differentiation are requisite processes for establishing cultivated crops. These processes inherently involve substantial changes in population structure, including those from artificial selection of key genes. In this study, accessions of peach (Prunus persica) and its wild relatives were analysed genome-wide to identify changes in genetic structures and gene selections associated with their differentiation. Analysis of genome-wide informative single-nucleotide polymorphism loci revealed distinct changes in genetic structures and delineations among domesticated peach and its wild relatives and among peach landraces and modern fruit (F) and modern ornamental (O-A) cultivars. Indications of distinct changes in linkage disequilibrium extension/decay and of strong population bottlenecks or inbreeding were identified. Site frequency spectrum- and extended haplotype homozygosity-based evaluation of genome-wide genetic diversities supported selective sweeps distinguishing the domesticated peach from its wild relatives and each F/O-A cluster from the landrace clusters. The regions with strong selective sweeps harboured promising candidates for genes subjected to selection. Further sequence-based evaluation further defined the candidates and revealed their characteristics. All results suggest opportunities for identifying critical genes associated with each differentiation by analysing genome-wide genetic diversity in currently established populations. This approach obviates the special development of genetic populations, which is particularly difficult for long-lived tree crops.

Key words: artificial selection, cultivar differentiation, domestication, linkage disequilibrium, tree crop

1. Introduction
Plant domestication establishes a vital co-dependence between humans and plants.1 Previous studies of the plant domestication process have focused mainly on seed-propagated annual (herbaceous) species, often with the goal of identifying wild progenitors, as well as changes in genetic structures associated with domestication.2–5 The results have provided historical insights into the genetic adaptation required for domestication, which has relevance to crop breeding strategies. During the domestication and early breeding process, crops typically experience population bottlenecks, including extensive artificial selection for improved crop quality and local adaptation.6 Evidence of this selection remains in the patterns of genetic diversity within cultivated...
Evolution of domesticated peach

2. Materials and methods

2.1. Sample collection and genome-wide genotyping

For genome-wide genotyping, leaves were collected from 67 accessions of domesticated peach (P. persica; subg. Amygdalus), 20 accessions of peach relatives (four Prunus davidiana, one P. kansuensis, 12 P. mira, one P. tangutica, and two P. webbii; subg. Amygdalus), and 8 accessions of outgroup species [two Japanese plum (P. salicina; subg. Prunus), two Japanese apricot (P. mume; subgen. Prunus), and four sweet cherry (P. avium; subg. Cerasus)], from the UC Davis and USDA Prunus germplasm collections in Winters, California, USA, the NIFTS Prunus germplasm collections, Tsukuba, Japan, the Research Institute for Agriculture Okayama Prefectural Technology Center for Agriculture, Forestry, and Fisheries, Akaiza, Japan, and the experimental orchard of Kyoto University, Kyoto, Japan, as summarized in Supplementary Table S1. Tissues from a total of 95 accessions were subjected to DNA extraction using Nucleon PhytoPure (GE Healthcare, Tokyo, Japan) and thereafter phenol/chloroform extraction.

Genotype calling was performed with an Illumina infinium peach 9K SNP chip, which was defined from a total of 1,022,354 SNPs that were identified from the resequencing data in a wide variety of 56 accessions in Amygdalus, mainly including peach (P. persica) and almond (P. dulcis), and ~75% of genic SNPs were verified in the peach genome.43,44 We assayed 7,873 SNPs in this study. Each 5,180 and 6,605 informative SNP was selected for structural analysis and evaluation of LD and selective sweeps, respectively (see below).

2.2. Allele pruning

SNPs showing >5% missing data or <0.05 minor allele frequency (MAF) in domesticated peach cultivars were pruned with PLINK.45 After filtering, 6,605 SNPs remained for use in estimating LD. SNP pairs showing strong LD were further pruned by defining a window of 50 SNPs, removing one of a pair of SNPs if $R^2 > 0.5$ (VIF threshold values = 2), and then shifting the window by three SNPs and repeating the procedure using PLINK. After filtering, 5,180 SNPs remained for use in the analyses of population structure.

2.3. Analysis of population structure

For the topology of the evolutionary tree, we aligned 5,180 concatenated SNPs to give two aligned positions for each SNP locus to reflect the diploid allelic states. The aligned concatenated sequences were
subjected to neighbour joining (NJ) using version 5.05 under the Poisson matrix with gamma distribution for the rates and with 1,000 bootstrap replications. Principal component analysis (PCA) was performed using the same 5,180 SNP set, using prcomp implemented in R version 2.15.3. Heterozygosity [expected (H\textsubscript{e}) and observed (H\textsubscript{o})], inbreeding coefficient (F\textsubscript{IS}), and Weir & Cockerham F statistics (F\textsubscript{ST}) were calculated with GENEOPOP 4.0.46,47 Identity by descent (IBD) proportions were calculated as pi-hat values with PLINK, considering all pairings of the 67 accessions of domesticated peach (P. persica). We used the pruned 5,180 SNPs for the calculation. We had no calibration to infer first- or second-degree relationships from the information of actual pedigrees owing to the lack of information on reliable pedigrees and to the small number of accessions.

To evaluate delimitations in population structure, we performed individual-based Bayesian clustering with Markov chain Monte Carlo (MCMC) simulations, using STRUCTURE 2.2.48 and InStruct49 to infer the population ancestry of genotypes in K predefined clusters. K values ranging from 2 to 10 were evaluated for subdivision of the full (domesticated or wild) peach population (n = 87). We fixed K at 2 for characterizing proportions of ancestry from two predefined ancestral gene pools in some combinations of the varietal complexes or of the resulting clusters, to support population subdivision, and to infer genetic introgression among the complexes according to Cornille et al.52 We performed an independent test of each K using at least 200,000 MCMC iterations after 50,000 burn-in iterations. To evaluate inference of K, the model with the highest ln Pr(K)48 and the ΔK model with the greatest second-order rate of change in ln Pr(K)48 were examined.

Pairwise LD among the SNPs were calculated with PLINK, using 6,605 SNP sets considering MAF (>0.05) and missing reads (>20%) in the full population as candidates. We exploited all R\textsuperscript{2} values in all pairs of SNPs in 10,000-kb windows. The R\textsuperscript{2} values were independent calculated for each subpopulation for each chromosome. The LD decay distances in each domesticated peach subpopulation, in the whole domesticated peach population, and in a wild related population were defined as the first points at which the average R\textsuperscript{2} values in a 100-kb bin revealed no significant difference (P > 0.1) against the background R\textsuperscript{2} values, which were calculated from average intra-chromosomal comparisons.

2.4. Haplotype phasing
We characterized haplotypes of SNPs in each chromosome with fastPHASE v.1.2 using 6,605 SNP sets considering MAF and missing rates in the entire population.59 In advance and against the full population, including 87 wild or domesticated accessions, we estimated imputation error rates using 1,583 SNPs on chromosome 4 and the following options in fastPHASE: number of random starts of the EM algorithm = 10, EM iterations = 25, lower limit numbers of clusters = 1, upper limit numbers of clusters = 20, and interval between values for number of clusters = 1. This analysis gave an imputation error rate of 0.0822 with K = 8 as an optimal condition, only slightly larger than the values reported for phasing of SNPs on grapevine chromosome 8 and in humans.18,53 We adopted K = 8 and constructed haplotypes for each chromosome. Construction of haplotypes was also performed using each subpopulation defined in structure analysis (see Results section), which often showed a lower imputation error rate (ca. 0.06–0.17) than the full population.

2.5. Detection of selective sweeps
The 6,605 SNPs considering MAF and missing rates were used for three approaches, site frequency spectrum (SFS)-, integrated Haplotype Score (iHS)-, and XP-extended haplotype homozygosity (EHH)-based methods. The SFS-based approach was applied using the pooled heterozygosity (H\textsubscript{p})59 of each cluster in a 400-kb sliding window with a 100-kb step. Windows contained only four or fewer positions where SNPs were removed. All SNP alleles were completely given in bi-allelic states, and thus, the pooled heterozygosity was given by the average expected heterozygosity in each window, as follows:

$$H_p = \frac{1}{S} \sum_{m=1}^{S} \left( \frac{2N(m)_{\text{maj}}N(m)_{\text{min}}}{(N(m)_{\text{maj}} + N(m)_{\text{min}})^2} \right)$$

Here, S is the number of SNP positions in a window, N(m)\text{maj} and N(m)\text{min} are the numbers of major and minor alleles, respectively, in mth SNP locus in a window. Individual H\textsubscript{p} values were then Z-transformed as follows: $ZH_p = (\bar{H}_p - \mu_{H_p})/\sigma_{H_p}$, according to a report on pooled heterozygosity for selective sweep analysis in chicken.59 The allele information was used in an unphased state.

Both iHS and XP-EHH tests were based on EHH\textsubscript{4}, which is defined as the probability that two randomly chosen chromosomes carrying the core haplotype of interest are identical by descent (as assayed by homozygosity at all SNPs) for the entire interval from the core region to point x.52 We detected EHH in each subpopulation using the program Sweep.5 We approached the computation of the integral of observed EHH (iHH)\textsubscript{4} in 1,000 kb from each SNP core by measuring EHH in every 100-kb bin. EHH decayed to below 0.05 by 1,000 kb from SNP cores in most cases.

For detection of iHS, iHH values from two core alleles at one SNP core position in one population were defined as iHHA and iHHD, which originally corresponded to ancestral and derived alleles, respectively.54 iHS statistic is then given as unstandardized iHS = ln(iHHA*iHHD) (1).54 Finally, we obtained the standardized iHS as a Z-transformed value of the unstandardized iHS. We examined the reciprocal states of two core alleles (iHHA and iHHD) because of the sparseness of information on the derivation of the core alleles among peach subpopulations. XP-EHH is also obtained by Equation (1), focusing on the same core alleles in comparison to two populations.5 For two populations, A and B, the log values of the integral EHH\textsubscript{A}, I\textsubscript{A} and I\textsubscript{B} (like iHHA and iHHD in the iHS test), ln(I\textsubscript{A}/I\textsubscript{B}) give an index of selection specific to either of the two populations. An unusually positive value of ln(I\textsubscript{A}/I\textsubscript{B}) suggests selection in population A, whereas a negative value suggests selection in population B. The standardized XP-EHH is given as a Z-transformed value of ln(I\textsubscript{A}/I\textsubscript{B}), the details of which are presented by Sabeti.53 The standardized iHS and XP-EHH were transformed to P values using R for graphical plotting.

2.6. Sequencing and genetic diversity analysis in selected regions
To exploit genes under selection during cultivar differentiation, full lengths of a total of 15 genes located on the region under putative selection were amplified by PCR using 30 peach accessions (Supplementary Table S3a) and then sequenced with an Illumina HiSeq 2000 (Illumina) as paired-end 100 PE100). The libraries were constructed based on in-house-developed protocols described previously.55 Approximately 4% of a sequencing lane was dedicated to all samples to yield at least 100x coverage in any region. Over 15% of read coverage for each SNP was considered as informative. All bioinformatic and statistical analyses were performed on local servers at the UC Davis Genome Center (Davis, CA, USA). Raw reads without adapter sequences were subjected to trimming (length > 35 bp, mean sliding window of 5 bp phred quality score ≥20) using custom Python scripts. Reads were then mapped to the reference sequences from the peach

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3. Results
3.1. Genotyping and defining the genetic structure
The topology of the evolutionary tree, constructed using 5,180 concatenated SNPs, showed distinct differentiation between peach and its wild relatives and also among the cultivar complexes, based on classical classifications by morphology or use (Fig. 1). Six major clusters were defined in the evolutionary tree: wild species (W), landraces (L), modern fruit cultivars (F), and modern ornamental cultivars (O-A and O-B for only ornamental usage and FO for fruit and ornamentals). The F (including FO) and O-A clearly diverged from landraces with strong statistical support (with bootstrap values of 70/100 and 74/100, respectively). The L clade was divided into two clades, of which one contained mainly East Asian cultivars (L-EA), and the other (L-OT) showed no specificity for geographic distribution. Some ornamental cultivars (O-B) were included in clade L-EA. Note that two wild species (P. davidiana 2325-21A, and P. mira 2228-21A) were located near or in landrace clusters, probably because of recent frequent hybridization with domesticated peach cultivars or ancestral shared polymorphisms, as supported by PCA and STRUCTURE analyses (see below).

To examine the structure in more detail, PCA, population structure, and $F_{ST}$ analysis were conducted for 87 Amygdalus subgenus
accessions using the 5,180 SNP set. PCA analysis showed distinct differentiation of the W clusters from *P. persica* with two exceptions (*P. davidiana* 2325-21A and *P. mira* 2228-21A) (Fig. 2A for the first two principal components). In addition, this analysis revealed that the F, FO, and O-A clusters had genetic structures that were distinct from those of the others, although O-A may have experienced some hybridization with the landraces in East Asia or ancestrally shared polymorphisms with them (Fig. 2A). The results of population structure analysis by STRUCTURE 2.2 and InStruct, assuming subpopulation \( K = 2 \)–10 distinct clusters, were consistent with each other, and supported almost the same conclusion as in the evolutionary tree and the PCA (Fig. 2B and C, for \( K = 2 \)–7). Model selection based on the \( \Delta K \) and \( \ln \text{Pr}(K) \) supported \( K = 6 \) (Supplementary Fig. S1A and B). With \( K = 3 \) or less, ornamental cultivars and landraces were not clearly separated, whereas with \( K = 7 \), they displayed different population structures, although some genetic hybridization was likely (Fig. 2B and C, Supplementary Fig. S1C for comparison of the O and L-EA clusters in predefining two ancestral gene pools). The FO cluster was inferred to have experienced clear hybridization with the F and O/L-EA clusters in \( K = 4 \)–7, a reasonable inference, given that they originated by recent hybridization between fruit and ornamental cultivars (Supplementary Fig. S1D). Two accessions of wild species, *P. davidiana* 2325-21A and *P. mira* 2228-21A, appeared to have experienced recent hybridization or ancestrally shared polymorphic alleles with landraces at \( K = 7 \), and for \( K = 2 \), their structures were almost the same as that of the domesticated *P. persica* (Fig. 2B and C). The \( F_{ST} \) test supported frequent hybridization in the FO cluster and the F and L-EA clusters and suggested significant differentiation in the domesticated peach (*P. persica*) from wild relatives (\( F_{ST} = 0.268 \) and 0.275 from *P. mira* and *P. davidiana*, respectively), and in F and O-A (\( F_{ST} = 0.158 \) and 0.135, respectively) from the landrace L-EA (Table 1), relative to the \( F_{ST} \) values for other tree crops and wild relatives.\(^{18,22}\) Also, the O-A and O-B clusters, both of which are categorized as ornamental varieties, showed distinct differentiation (\( F_{ST} = 0.228 \)).

Based on these population structure results, the O-B cluster was included in the L-EA (or L) cluster and the F-O was no longer considered because at least one of them originated from the recent hybridization between F and O-A gene pools (Supplementary Fig. S1D).

**Figure 2.** Population structure analysis in peach. (A) Principal component analysis using information from 5,180 genome-wide SNPs in 87 accessions including domesticated peach and its wild relatives (right) and in 67 accessions focusing on domesticated peach (left). The first two components in PCA (PC1 and PC2) are plotted on the axes to visualize the genetic relationships. The proportion of variance explained by each PC is given in parentheses along each axis. The wild species is shown as a cross. The landraces are shown as squares in green and yellow for accessions in East Asia and other regions, respectively. The modern cultivars are shown as circles in blue and red, corresponding to fruit and ornamental cultivars, respectively. (B and C) Structure analysis of subdivision of the population \((K = 2-7)\), with STRUCTURE 2.2 (B) and with InStruct (C). Each individual is shown as a vertical bar. In \( K = 2 \), wild species showed a cluster distinct from domesticated peach, except for two accessions experiencing frequent hybridization with domesticated peach. In \( K = 5 \) or more, the F, L-OT, and O-A/O-B/L-EA clusters show clear separation. The O-A and O-B/L-EA showed significant separation at \( K = 7 \) or more.
were also supported by FIS values, suggesting degrees of inbreeding to the W cluster revealed no significant over, the FO cluster demonstrated significant LD in landraces such as grape (up to 50–150 kb),9,60 sorghum (up to ca. 150 kb),42 and maize (up to 2 kb).41 In comparing clusters, the F cluster showed much greater delay in decay of LD than the two subclusters of the L cluster (L-EA and L-OT), ranging over 3,000 kb on some chromosomes (Fig. 3B, Supplementary Fig. S3). This finding supports a bottleneck or shift to higher inbreeding in the establishment of the F cluster from the progenitor landraces. The results of IBD test would support a recent bottleneck from cv. Chinese Cling (Supplementary Fig. S2). The O-A cluster, in general, showed no significant differences in LD decay from the two L subclusters (Fig. 3B), although a slightly significant increase in LD could be detected for the O-A cluster on a few chromosomes (Supplementary Fig. S3). A significant delay in LD decay might also be expected from the distribution of EHH presented later. In the genome-wide LD analysis, we detected a clear monotonic decrease in static values of LD with the physical distance on any chromosome ranging from at least 1,000 kb in the domesticated peach cluster to even more in the F subcluster. This situation would make it preferable to identify selective sweeps based on LD index using small numbers of marker sets. In this study, we could use at least 5,000 pruned SNP sets (one SNP locus per ca. 45 kb on average in the peach genome) for subsequent analysis of selected genomic regions, yielding sufficient information for the identification of statistically distorted LD values from each SNP core in the genome.

### 3.3. Genome-wide detection of selection in domestication and cultivar differentiation pathways

To evaluate selective sweeps in local genomic regions, indexes based on the site frequency spectrum (SFS), such as \( \pi \) or Tajima’s \( D \) which can detect fixed sweeps in a population, have been applied in plant species.8,12,14,30 In genome-wide analysis of selected regions, similar approaches including LD- or haploblock-based methods, which usually detect very recent and ongoing sweeps, are reported to be effective for some plant species8,10,62 and for some domesticated animals such as dog63.
peach genome, distortion of the availability in SNP patterns, or simple drift cannot be ruled out. Comparison of the \( H_\text{p} \) values between the F- or O-A clusters and the L cluster (Fig. 4B) revealed a significant bias in homozygous states among the clusters in specific genomic regions and suggested that at least some represent true selective sweeps that specifically occurred in the path to modern fruit or ornamental cultivar development. The middle of Chromosome 7 (ca. 8,000–8,800 kb) and the bottom of Chromosome 4 (ca. 29,200–29,600 kb) showed particularly clear tendencies to selective sweeps specific to the F and O-A subpopulations, respectively (Fig. 4C and D).

LD- or haploblock-based analyses were performed in accordance with the concept of EHH.\(^{52}\) The program Sweep\(^{53}\) was used to evaluate the \( i\text{HS} \),\(^{54}\) which quantifies the difference of EHH values around the selected locus in one population, and the Cross-Population EHH (XP-EHH),\(^{55}\) which calculates EHH values from the same SNPs core between two populations using phased SNP strings. In both analyses, EHH values collected in 1,000-kb sections from each SNP locus showed significant reductions (ca. 0.05).

The \( i\text{HS} \) in domesticated peach, as well as the two subpopulations F and O-A, showed specific patterns of significant peaks (\( P < 0.0001 \)) corresponding to putative selective sweeps (Fig. 5A, Supplementary Figs S5 and S6A). The patterns of the peaks for the selective sweep in \( i\text{HS} \) were considerably different from those in the SFS-based analysis (Fig. 4), perhaps because the \( i\text{HS} \) detects mainly selected alleles in the heterozygous state, whereas the SFS-based analysis is applicable mainly to the detection of homozygosity. Still, the same positions for putative selection in SFS-based and \( i\text{HS} \) analyses were selected on Chromosome 7 (ca. 8,000–8,800 kb) in the F cluster (\( P > 0.00001 \) in \( i\text{HS} \)). The XP-EHH analyses were performed in comparisons between domesticated peach and the W clusters, and between the F or O-A and the L clusters (Fig. 5B and Supplementary Fig. S6B) to analyse genes selected in the course of domestication and cultivar differentiation. In the XP-EHH analysis of domesticated peach and the W clusters, the peaks were at the bottoms of Chromosome 4 (ca. 25,000–27,000 kb for \( P < 0.001 \)), and Chromosome 6 (ca. 25,000–26,500 kb for \( P < 0.001 \)). In XP-EHH analyses, they were similar to those in \( i\text{HS} \) analyses of the domesticated peach (Fig. 5, \( P < 0.00001 \) for chr. 4, and \( P < 0.0001 \) for chr. 6). Some of the significant peaks in XP-EHH analyses were different from those in the \( i\text{HS} \) and SFS-based approaches. Similar situations have been reported in analyses using EHH- and SFS-based methods in human evolution, which captured certain selected regions in all approaches, further supporting the value of using multiple approaches for comprehensive analysis of positive selection.\(^{52,53}\) The XP-EHH analyses of the F or O-A cluster relative to the L cluster also showed some specific peaks, though different from the \( i\text{HS} \) and SFS-based results. Significant peaks could still be detected on Chromosome 4 (ca. 8,500–9,500 kb) and Chromosome 7 (ca. 8,000–8,800 kb) in the F cluster (\( P < 0.00001 \)). The peak on Chromosome 4 was also detected in the domesticated peach cluster by the SFS-based method (Fig. 4).

### 3.4. Candidate genes under selection

For the F and O-A clusters, genetic diversity of candidate genes located on representative regions under putative strong positive selection was estimated. These included Chromosome 7 (8,000–8,800 kb) for the F cluster, Chromosomes 4 (29,200–29,400 kb) and 8 (16,900–17,550 kb) for the O-A cluster, and Chromosome 1 for both (1,200–1,500 kb), using the 30 domesticated peach accessions (Supplementary Table S3). Note that the focus was only on genes annotated with functions possibly conferring advantages of differentiation. Of these, ppa004528m, ppa016246m, and ppa025156m showed significant reduction in genetic diversity in the F cluster (\( \pi = 0.00007 \)) against the O-A and

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**Figure 3.** Comparison of LD decay in the wild species and each variety complex of domesticated peach. The average values of \( R^2 \) in pairwise LD among the 1,346 SNPs on Chromosome 4, which carries the most SNPs among the eight chromosomes, in windows of up to 10,000 kb are shown. The Y-axis standards are adjusted according to the background values in each subpopulation for visualization of LD decay for comparison among subpopulations. The X-axis shows physical distance among the SNPs, and the average \( R^2 \) values at 100-kb intervals are plotted. (A) Comparison of LD decay in domesticated peach and wild relatives. Black and white arrows indicate LD decay points in domesticated peach and wild relative, respectively. (B) Comparison of LD decay in the F, O-A, L-EA, and L-OT clusters. The F cluster shows only significantly expanded LD, as shown by the black arrow, in comparison to the other clusters, whose LD decays are shown with white arrows and black bars.

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and chicken.\(^{39}\) Here, selections for domesticated peaches were expected to be recent or ongoing, particularly for the F cluster. Peach is a species with a long generation time (at least 2–3 yrs to flowering) and one for which individual genotypes can be maintained by vegetative propagation for long time periods, suggesting that selected genes may have been maintained in heterozygous states. Thus, we adopted indexes based not only on SFS but also on LD or haploblock for exploiting selective sweep, following genome-wide analyses of selection in human genomes.\(^{73–54}\)

In SFS-based analysis, we evaluated the transition of pooled heterozygosity (\( H_\text{p} \)) of each cluster using informative SNPs in a 400-kb sliding window with 100-kb steps, following a previous report on SFS-based analysis in chicken.\(^{39}\) The distributions of observed \( H_\text{p} \) values and Z-score values of \( H_\text{p} \) (\( ZH_\text{p} \)), which considered coalescent effects by using the entire SNP sets in the peach genome, are shown in Fig. 4A for the domesticated peach population and two subpopulations, F and O-A. All showed some genomic regions containing candidates for selective sweeps with significant homozygous states in comparison to the whole genome (\( P < 0.001 \)). However, the possibility that they are derived from differences in the substitution ratio in the
Figure 4. Genome-wide selective sweep analysis based on $ZH_p$ in the domesticated peach and two modern varietal complexes. (A) Transitions of $Z$-transformed values of $H_p$ in 400-kb bins with 100-kb steps are shown for the domesticated peach population and two subpopulations, the F and O-A clusters. Putative regions showing selective sweeps ($P < 0.001$) are indicated by outlined triangles. (B) For the F and O-A clusters, plots of relative values of $H_p$ in 400-kb bins with 100-kb steps against the L cluster are shown. Putative regions showing selective sweeps in the paths of differentiation from landraces are indicated by triangles. For the two regions indicated by black triangles, detailed characterization of $ZH_p$ in a comparison among four subpopulations (F, O-A, L-EA, and L-OT) is shown in (C) for the top of Chromosome 7, which corresponded to a putative selective sweep in the F cluster, and in (D) for the bottom of Chromosome 4, showing a putative selective sweep in the O-A cluster.
landrace (L-EA and L-OT) clusters (average $\pi = 0.00114 \pm 0.00021$, $0.00279 \pm 0.00011$, and $0.00629 \pm 0.00091$, respectively, Supplementary Table S4). However, no disruptions or substitutions clearly associated with the functional changes in the ORF sequences could be identified, although minor substitutions on protein sequences were observed in comparison to some landraces. For the O-A cluster, ppa003808m showed a significant reduction in genetic diversity ($\pi = 0$) in comparison to the other clusters (average $\pi = 0.00150 \pm 0.00017$, Supplementary Table S4). ppa003808m retained a trihelix motif, showing high similarity to the PETALLOSS (PTL) gene, which is involved in petal size and morphology in Arabidopsis thaliana.64–66

4. Discussion
4.1. Domestication path in peach
The domestication events for major annual crops, such as maize or rice, have been well characterized, owing in part to the relatively straightforward selection for consumable parts as well as the genetic fixation of key controlling genes in distinct species or cultivars. Tree crops appear to have more complicated domestication histories.18,22 For example, landrace accessions of P. persica, such as Kemomo Nagoshijou or Kutao, which show wild species-like characteristics, share a similar genetic structure with cultivated peach accessions. Thus, it
may be difficult to define a single domestication path. Two scenarios are supported by the data: (i) current wild peach cultivars (fetal peaches) were derived from divergent lines after a general domestication event or (ii) multiple independent domestications occurred for each trait or region. The first scenario (i) would fit well with the premise that P. persica was first domesticated from a wild species in China before diffusion to other regions.28,29 Such a general domestication event is supported by the tendencies for population bottlenecks or inbreeding to be detected using the full population of peach (P. persica) in this study. In contrast, other domesticated perennial crops such as grape and apple are reported to lack narrow domestication bottlenecks.18,22,67 However, in theory, perennial woody crops are under strong bottleneck pressures during domestication, resulting from the ease of their clonal propagation as well as difficulties associated with their long generation time. The wide domestication bottleneck in grape might be explained by widespread use of vegetative propagation of a large number of landraces with a resulting maintenance of genetic diversity as well as of many local genetic clusters.18 This situation would be more aligned with the second scenario (ii). The results of IBD analysis in peach showed that, except for the F cluster, most cultivars have no first- or second-degree relationships. This situation is distinct from that of domesticated grapevine, which shows a high ratio of first-degree relationships with a relatively small number of representative cultivars.18 This finding indicates that there were not many opportunities for domestication or construction of local clusters in peach, at least for the cultivars examined in this study.

4.2. Selection in peach domestication and cultivar differentiation

Ancient selection in the peach genome appears likely, although some observations may be due to genetic drift. Results from QTL analysis have also been used to support selection during domestication and cultivar differentiation in rice8 and chicken15. In the present study, the strong putative selection on Chromosome 4 (ca 8,500–9,500 kb) detected in modern fruit cultivars (the F cluster) is consistent with a major QTL (UDP96–003) for important domestication traits, including fruit size and maturity date previously reported by Qiuot et al. in a backcross between cultivated peach and its wild relative P. davidiana.68 This finding suggests that favoursable allele(s) in this region have been selected from wild ancestors during domestication. Furthermore, evidence for selection is observed at the bottom of Chromosome 6 (ca 2.5,000–26,500 kb) which corresponds to the genomic regions XP-EHH and iHS, known to contain the self-incompatibility (S) controlling haplotype, including an F-box gene as the pollen-S (SFB) factor and an RNase gene as the pistil-S (S-RNase) factor in Prunus species.69,70 Previously, it has been suggested that the whole domesticated peach (P. persica), including all clusters apart from the W cluster in this study, is a self-compatible (SC) species with at least four forms of pollen-S, PpSFB1-4, disruptive mutations.71,72 Meanwhile, to some extent, the wild species (W cluster) in the Amygdalus subgenus are supposed to demonstrate SI.73 In general, a shift from SI to SC is strongly selected during domestication to facilitate the stable production of inbred lines.27 In contrast, a change in the mating system from SI to SC could have a strong influence on the pattern of polymorphisms, affecting genetic diversity and LD. This might result in the differences in genetic structures between wild species and domesticated peach.

At least three candidates for selection were identified in the F cluster, with many on Chromosome 7. The candidates, ppa004528m, ppa016246m, and ppa025156m, show high similarity to lycopene cyclase-like At3g10230 (LYC/lAt5g57030 (LUT2), At3g18030 (Arabidopsis thaliana Hal3-like protein A; AtHAL3A), and At1g18990 (reduced vernalization response 1; VRN1) in the Arabidopsis thaliana genome. Lycopene cyclases (LYCs) play a role in the biosynthesis of lutein, which is a member of the carotenoid pathway.74 The simplest explanation for the selection of ppa004528m may be its association with the yellow carotenoid pigmentation in peach. However, the flesh coloration of peach is determined mainly by the Y locus, which is reportedly located on Chromosome 1 (LG1).75,76 Major QTLs for skin color are also different from those in most regions identified as having experienced selective sweeps, although one of them is located on Chromosome 7.77 The other two candidates, AtHal3A-like and VRN1-like genes, may contribute salt/osmotic tolerance and flowering/bud-burst timing, respectively, based on the functions of their homologs in Arabidopsis.78,79 In considering the uses and characteristics of modern fruit cultivars, we may expect that some genes directly involved in fruit traits, such as fruit size, sugar contents, or the important flesh-softening trait often called ‘melting texture’, have been under strong selection. Where selection for more ornamental traits is expected, as in the O-A cluster, ppa003808m appears to be a good candidate, given that the associated PETALLOSS trait is known to affect morphological or architectural changes, particularly for flowers.

Genes with annotation for fruit quality and ornamental value are limited at present, presumably because of the lack of information for these traits in model plant species. Thus, the still-uncharacterized genes located in the selected regions in the F and O-A clusters may provide opportunities for elucidating critical tree domestication factors not present in current annual plant models. Genes controlling such key domestication and cultivar differentiation factors could be identified by improved characterization of the selection pathways involved, perhaps by further exploiting the wide genetic diversity present in peach and similar perennial tree crops. This research approach could prove powerful, particularly for long-lived perennial crops where mutagentic and map-based approaches are restricted.

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Supplementary data

Supplementary data are available at www.dnaresearch.oxfordjournals.org.

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