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Phenotypic and fine genetic characterization of the D locus controlling fruit acidity in peach

Karima Boudehri¹, Abdelhafid Bendahmane², Gaëlle Cardinet¹, Christelle Troadec², Annick Moing³,⁴ and Elisabeth Dirlewanger*¹

Address: ¹INRA, UR0419, Unité de Recherches sur les Espèces Fruitières, Centre de Bordeaux, BP 81, F-33140 Villenave d’Ornon, France, ²INRA-CNRS, UMR1165 Unité de Recherche en Génomique Végétale (URGV), 2 rue Gaston Crémiex, F-91057 Evry, France, ³INRA – UMR619 Fruit Biology, INRA, Université de Bordeaux 1, Université de Bordeaux 2, BP 81, F-33140 Villenave d’Ornon, France and ⁴Metabolome-Fluxome Pole, IFR103 BVI, BP 81, F-33140 Villenave d’Ornon, France

Email: Karima Boudehri - kboudehr@bordeaux.inra.fr; Abdelhafid Bendahmane - bendahm@evry.inra.fr; Gaëlle Cardinet - gaelle.cardinet@bordeaux.inra.fr; Christelle Troadec - troadec@evry.inra.fr; Annick Moing - moing@bordeaux.inra.fr; Elisabeth Dirlewanger* - dirlew@bordeaux.inra.fr

* Corresponding author

Abstract

Background: Acidity is an essential component of the organoleptic quality of fleshy fruits. However, in these fruits, the physiological and molecular mechanisms that control fruit acidity remain unclear. In peach the D locus controls fruit acidity; low-acidity is determined by the dominant allele. Using a peach progeny of 208 F₂ trees, the D locus was mapped to the proximal end of linkage group 5 and co-localized with major QTLs involved in the control of fruit pH, titratable acidity and organic acid concentration and small QTLs for sugar concentration. To investigate the molecular basis of fruit acidity in peach we initiated the map-based cloning of the D locus.

Results: In order to generate a high-resolution linkage map in the vicinity of the D locus, 1,024 AFLP primer combinations were screened using DNA of bulked acid and low-acid segregants. We also screened a segregating population of 1,718 individuals for chromosomal recombination events linked to the D locus and identified 308 individuals with recombination events close to D. Using these recombinant individuals we delimited the D locus to a genetic interval of 0.4 cM. We also constructed a peach BAC library of 52,000 clones with a mean insert size of 90 kb. The screening of the BAC library with markers tightly linked to D locus indicated that 1 cM corresponds to 250 kb at the vicinity of the D locus.

Conclusion: In the present work we presented the first high-resolution genetic map of D locus in peach. We also constructed a peach BAC library of approximately 15x genome equivalent. This fine genetic and physical characterization of the D locus is the first step towards the isolation of the gene(s) underlying fruit acidity in peach.
Background

Peach [Prunus persica (L.) Batsch] belongs to the Spiraeoideae subfamily of the Rosaceae [1]. The Prunus genus is characterized by species producing drupes as fruit, and contains a significant number of economically important fruit tree species such as almond (Prunus dulcis (Mill.)), apricot (Prunus armeniaca L.), sweet cherry (Prunus avium L.), sour cherry (Prunus cerasus L.) and plum (Prunus domestica L.).

Compared to other tree species, peach has a relative small diploid genome (290 Mb) [2], and a short juvenile phase (two to three years). Therefore, peach is considered as a model species for Rosaceae family and a physical map of its genome has been initiated [3].

Among fruit producing rosaceous crops, peach is the second most important fruit crop in Europe after apple and the third worldwide (FAOSTAT: http://faostat.fao.org/). However, the consumption of peaches and nectarines is stagnant due to the low quality of fruits that are harvested at an immature stage for storage and shipment reasons [4]. One of the major objectives for peach breeders is to find the right compromise between quality and immaturity at harvest [5]. The variation in fruit quality at harvest involves a large number of interrelated factors [6] among which organic acid and soluble sugar contents and composition are major determinants [7]. In ripe peach fruit, malic and citric acids are the predominant organic acids, while quinic acid accumulates in lower amounts [8,9]. Moreover, the major soluble sugars are sucrose, fructose, glucose and sorbitol [9,10]. Sucrose is the predominant soluble sugar at maturity while sorbitol accumulates at very low levels.

In peach, the D locus (D is for 'Doux' meaning 'sweet' in French) was described as dominant and controlling the 'low-acid' character of fruit [11,12]. Based on previous segregation analyses of an F2 population (JxF) obtained from a cross between 'Ferjalou Jalousia'™ low-acid variety and 'Fantasia' normally-acid variety, the D locus was mapped on linkage group 5 [13]. It is co-localized with major QTLs for pH, titratable acidity (TA), organic acids concentration and with small QTLs for sugars concentration [14]. Low-acid peach fruit is characterized by reduced contents of malic and citric acids [9], which, however, cannot be explained just by the reduced expression or activity of phosphoenolpyruvate carboxylase (PEPC) [15], a key enzyme involved in malate synthesis. 'Ferjalou Jalousia’™ fruit has half the concentration of malic acid and one-fifth that of citric acid of 'Fantasia' variety [9]. Using the candidate gene approach, 18 genes involved in organic acid synthesis, degradation or vacuolar storage were studied [16,17]. Expression analyses in fruit of six selected candidate genes did not show a clear difference between the normally-acid and low-acid varieties [17]. The genes showing a modification of their expression in the low-acid fruit compared to the normally-acid fruit were the tonoplastic proton pumps PRUpe;AtpvA1, PRUpe;Vp2, and to a lesser extent PRUpe;Vp1. PRUpe;Vp1 and PRUpe;Vp2 at citric acid peak and maturity, and PRUpe;AtpvA1 during cell division showed higher expression in the fruit of the low-acid variety ('Ferjalou Jalousia™'). However, none of these candidate genes were located on linkage group 5, excluding their direct role in the control of acid content by the D locus [17]. More recently, in the European ISAFRUIT Integrated Project http://www.isafruit.org/Portal/index.php, several candidate genes involved in fruit quality were selected and tested on the JxF F2 mapping population. However, none of them was located in the region of the D locus (Dirlewanger E., manuscript in preparation).

Low-acid varieties have already been described in apple [18], tomato [19], grape [20] and several Citrus species [21]. In apple, a non-acid mutant from the 'Usterapfel' variety showed a content in malic acid ten times less than the normally-acid one [22].

The high level of malic acid was reported to be controlled by the dominant Ma allele [23] suggesting that Ma and D act at different physiological control points. A cDNA-AFLP analysis, coupled with a bulk segregant analysis (BSA) was recently used to screen genes differently expressed between low- and high-acid apple fruits [24]. The authors reported the isolation of a cDNA whose expression could only be detected in low-acid fruit at an early stage of fruit development. Nevertheless, this cDNA showed no homology with any sequences in public databases. Moreover, the Ma and D loci are not located on homologous chromosomes: Ma is located on linkage group 16 in Malus [25], homologous to linkage group 1 in Prunus [26] and D is located on linkage group 5 in Prunus which is homologous to linkage groups 6 and 14 in Malus [27]. For Citrus, the low level of citric acid is controlled by a recessive gene named acitric [28]. Fruit acidity in Citrus seems to be linked to the capacity to accumulate citric acid into the vacuole. Low-acid varieties accumulate low amount of citric acid probably because it is exported out from the vacuole [29,30]. Two candidate genes such as acid invertase and cytoplasmic isocitrate dehydrogenase were identified to be differentially expressed between acid and low-acid Citrus [30]. Fruit acidity can also be controlled by several chromosome regions as in tomato where several QTLs for titratable acidity and pH were identified [31,32] and several candidate genes were proposed [33]. However, to date the mechanism(s) of the genetic control of fruit acidity remains to be elucidated.

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In order to identify genes of interest, candidate gene approach can be used when assumptions can be made regarding the biological function of the gene [34]. This approach was successfully undertaken for several fruit traits including anthocyanin content for which the biosynthesis pathway and regulating genes were well known [35] and cell wall degradation where implicated genes were identified in other species [36]. However, to isolate agronomically important and botanically relevant genes with unknown function and where no clear hypothesis can be made, chromosome landing seems the main strategy by which map-based or positional cloning could be applied [37]. The complexity of organic acids metabolic pathways as well as the difficult understanding of the regulation of their transporters and channels and related proton pumps [38,39] has hampered, so far, the identification of the gene(s) associated to the D locus using a candidate gene approach. Thus, in order to understand the molecular and physiological bases of this trait, a positional cloning strategy was initiated and a fine map of the D locus has been constructed. To identify the gene(s) underlying acidity control at the D locus, the first step was to construct a fine map of the D locus. The aims of the present work were: (1) the characterization of the fruit acidity trait, (2) the increase of the number of markers tightly linked to the D locus, (3) the conversion of the nearest markers into Sequence Characterized Amplified Region (SCAR) and Cleaved Amplified Polymorphic Sequence (CAPS) markers, (4) the construction of a high-resolution genetic map of this locus and definition of the position of the D locus with new recombinant individuals phenotyped, and (5) the evaluation of the genetic distance/physical distance ratio around the D locus using a BAC library.

Results

Fruit acidity characterization

Among the 208 individuals used for the genetic linkage map, only 151 trees producing fruit were phenotyped for pH and titratable acidity and were classified into three subgroups corresponding to the three genotypes: homozygous for 'Ferjalou Jalousia' allele (JJ) and for 'Fantasia' allele (FF), and heterozygous (JF) at the targeted locus (Fig. 1). A significant difference (Student's t-test, \( P < 0.01 \)) was observed for pH and TA for the comparisons of JJ and JF genotypes, FF and JF genotypes, and JJ and FF genotypes (pH mean values JJ = 4.57, JF = 4.36, FF = 3.63; TA mean values JJ = 36.5, JF = 48.2, FF = 109.7 meq/l) suggesting that the D allele is partially dominant. Homozygous JJ genotypes showed values higher than 4.12 for pH and lower than 51.9 meq/l for TA. On the opposite, pH and TA values for homozygous FF genotypes were respectively lower than 3.93 and higher than 65.6 meq/l. The pH of heterozygous JF genotypes ranged from 3.80 to 4.87 and TA ranged from 28.1 to 90.0 meq/l. Thus, normally-acid phenotypes that correspond exclusively to genotypes FF showed pH value lower than 3.8 and TA value higher than 100 meq/l while low-acid phenotypes corresponding to genotypes JJ or JF showed pH values higher than 4.0 and TA value lower than 60 meq/l. These results indicate that individuals with intermediate acidity (pH values between 3.8 and 4.0 and TA values between 60 and 100 meq/l) can be either homozygous dd or heterozygous Dd at the D locus and therefore, they cannot be reliably classified into normally-acid or low-acid phenotype.

Identification and mapping of AFLP markers linked to the D locus

Among the 1,024 primer combinations tested, 960 provided readable amplification products. Thirty to 90 bands were observed on AFLP gels per primer combination with a size range from 60 to 1,000 bp, but only 6.5% of the bands were polymorphic between the 'Ferjalou Jalousia' and 'Fantasia' parents. Markers whose bands were present in BDD1 and BDD2 bulks and absent from BDD1 and BDD2 bulks were potentially linked to the D locus (Fig. 2). A total of 34 markers were identified as putatively linked to the D locus (Table 1). Nineteen primer combinations each revealed only one D-linked marker, six primer combinations produced two D-linked markers (pGC-AGG, pCA-GCG, pTC-CAC, pCA-ACC, pCA-TCC and pAA-ACA) and one primer combination revealed three D-linked markers (pGC-TCT). As expected, all 34 AFLP markers were mapped on linkage group 5. AFLP markers close to the D locus (within 22 cm) were clearly polymorphic between bulks. AFLP markers mapped further away (beyond 27 cm) were polymorphic markers between bulks but with a very faint band for ‘d’ bulks. Fourteen AFLP markers were located within the first 10 cm containing the D locus.

Conversion of AFLP markers into SCAR and CAPS markers

Nine AFLP markers linked to the D locus were converted into simple codominant PCR-based markers. Four of them were codominant markers and five were dominant markers (Table 1). The codominant AFLP markers (pGC-AGG, pAC-AAC, pTC-CAC, pCA-ACC, pCA-TCC and pAA-ACA) were successfully converted into SCAR markers (D-Scar0, D-Scar1 and D-Scar2) and were confirmed as codominant markers (Table 2, Fig. 3). The codominant AFLP markers pTC-GTA, pGC-TCT revealed a deletion of one nucleotide in 'Fantasia' compared to 'Ferjalou Jalousia'. After sequencing the two alleles, three single nucleotide polymorphisms (SNPs) were detected; one of them was revealed after digestion with the restriction enzyme AciI and directly observed on agarose gel. This codominant Cleaved Amplified Polymorphism Sequence (CAPS) marker was named D-Scar3 (Fig. 3).
Figure 1
Frequency distribution of pH and titratable acidity of fruit juice. Mean values observed in 2007 are represented. Distribution of F₂ individuals, having fruit at maturity, used for the genetic linkage map. White, black and grey bars indicate homozygous genotypes for 'Fantasia' allele, 'Ferjalou Jalousia®' allele and heterozygous, respectively.
For the dominant AFLP markers, primers were designed from the sequences of individuals carrying the D allele. They were then tested on 'Ferjalou Jalousia®', 'Fantasia' and on the F 1-JF:21 hybrid used to construct the F 2 mapping progeny. The comparison of the sequences obtained for the pTG-TGG470J marker revealed a deletion of six nucleotides in 'Fantasia' as compared to 'Ferjalou Jalousia®' sequence. This AFLP marker was then transformed into a codominant SCAR marker (D-Scar6) (Fig. 3). For pTC-CTG470J, the sequencing of the two alleles revealed one SNP. The alleles can be discriminated by digesting the PCR products with MseI. This CAPS marker was called D-Scar7 (Fig. 3). For the three dominant AFLP markers pGT-TTG188J, pCA-GCG149F and pCA-GCG132J, no polymorphism was detected.

The six polymorphic SCAR markers were subsequently used to genotype the 208 individuals of the genetic linkage map; it confirmed that their localization was the same as AFLP markers (data not shown).

Furthermore, considering the total size of the obtained sequences (2,711 bp), this analysis revealed seven SNPs. Based on these results the frequency of the SNPs at the vicinity of the D locus was estimated to 2.6 SNPs per kb.

### High resolution mapping of the D locus

The fine mapping of the D locus was performed in two steps using the six SCAR markers described in the present study and three SSR markers MA026a, BPPCT041 and CPPCT040 already mapped to the proximal end of the linkage group 5 [13,26]. The first step was to genotype the 1,718 individuals from the seven segregating populations with three SCAR markers (D-Scar0, D-Scar2 and D-Scar6) and two SSR markers (MA026a and BPPCT041) spanning a region of 10.2 cM around the D locus (Fig. 4). A total of

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**Table 1: AFLP markers mapped on linkage group 5 (LG5) based on 208 JxF F2 individuals**

| AFLP marker | Position on LG5 (cM from the top) | Selection for conversion into SCAR | SCAR marker |
|-------------|----------------------------------|-----------------------------------|-------------|
| pAC-AAC402J | 0.7 | Yes | D-Scar1 |
| pGG-TAC215J | 0.7 | Yes | D-Scar2 |
| pGC-AGG430J | 0.7 | Yes | D-Scar0 |
| pTC-CTG470J | 1.8 | Yes | Monomorphic |
| pCA-GGA149F | 3.3 | Yes | D-Scar3 |
| pTC-CAC306 | 4.1 | Yes | Monomorphic |

* Allele sizes for both parents are indicated for codominant markers, while only allele size for one parent is indicated for dominant markers. 1 'Ferjalou Jalousia®', 2 'Fantasia'.

For the dominant AFLP markers, primers were designed from the sequences of individuals carrying the D allele. They were then tested on 'Ferjalou Jalousia®', 'Fantasia' and on the F1-JF:21 hybrid used to construct the F2 mapping progeny. The comparison of the sequences obtained for the pTG-TGG470J marker revealed a deletion of six nucleotides in 'Fantasia' as compared to 'Ferjalou Jalousia®' sequence. This AFLP marker was then transformed into a codominant SCAR marker (D-Scar6) (Fig. 3). For pTC-CTG470J, the sequencing of the two alleles revealed one SNP. The alleles can be discriminated by digesting the PCR products with MseI. This CAPS marker was called D-Scar7 (Fig. 3). For the three dominant AFLP markers pGT-TTG188J, pCA-GCG149F and pCA-GCG132J, no polymorphism was detected.

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308 individuals were found to have at least one recombination event between the farthest markers, MA026a and BPPCT041. The second step was to genotype the resulting 308 recombinant individuals with the three other SCAR markers (D-Scar1, D-Scar3 and D-Scar7) and CPPCT040 which were located within the spanned region. According to the recombination events between these nine markers, it was possible to determine the precise marker order on linkage group 5 (Fig. 4).

For the 149 recombinant individuals which produced fruits in 2007 and 2008, TA mean values varied from 15 to 167 meq/l and pH mean values ranged from 3.36 to 5.59 (Fig. 5). Among these recombinants, 110 individuals

Table 2: SCAR and CAPS markers developed from AFLP fragments linked to the D locus

| AFLP marker* in bp | SCAR marker | Primer sequence (5'-3') | Size¹ (bp) | P² | Annealing temp. (°C) | Enzyme |
|--------------------|-------------|-------------------------|------------|----|---------------------|--------|
| pGC-AGG<sub>430</sub>-<sub>450</sub>F | D-Scar0 | F GTGCACAGCTATCTCTTTTC, R CTCATGGCAACAAACATTC | 160 (J), 175 (F) | SSR | 52 | no |
| pAC-AAC<sub>402</sub>-<sub>412</sub>F | D-Scar1 | F GGGATGTGGGTATGTCGTA, R ACAAGGAGGAAGCTCTGG | 345 (J), 364 (F) | SSR | 55 | no |
| pGG-TAC<sub>215</sub>-<sub>221</sub>F | D-Scar2 | F CTCTACGTCTACGACGACAAC, R TGAGTCCTGAGTATTGTTGCTCTG | 142 (J), 148 (F) | Indel | 54 | no |
| pTC-GTA<sub>218F</sub>-<sub>219F</sub> | D-Scar3 | F GTTGAACATGAAACAAATGACATTG, R CAGTCGTCTCTCTTGAATCAGTCACATGC | 180 (J, F), 186 (F) | SNP | 52 | AccI |
| pTG-TGG<sub>470</sub>J | D-Scar6 | F CATGGCCCATCTCTTTTAC, R GACCAGTGGCATCTACTTCATTG | 92 (J), 98 (F) | Indel | 55 | no |
| pTC-CTG<sub>470</sub>J | D-Scar7 | F CTGCTCAGCTACCTGGTCTC, R TCCAACCTCAGGCTTGC | 334 (J, F), 340 (F) | SNP | 55 | MseI |

¹ (J) 'Ferjalou Jalousia®', (F) 'Fantasia'. ² Polymorphism observed.
were classified as producing low-acid fruit, 12 were identified as producing normally-acid fruit and 27 were considered as intermediate and were therefore, not classified. Among the individuals producing low-acid fruit, only 40 individuals recombining from heterozygous (Dd) to homozygous (dd) were informative. Then, only 52 recombinant individuals with extreme values for pH (from 3.36 to 3.68 for individuals with normally-acid fruit and from 4.20 to 5.55 for individuals with low-acid fruit) and TA (from 104 to 167 meq/l for individuals with normally-acid fruit and from 17 to 56 meq/l for individuals with low-acid fruit) were used to identify the position of the D locus. Among the 52 recombinant individuals so selected, 36 recombinant individuals between CPPCT040 and BPPCT041 indicated that the D locus was located upper than CPPCT040, while fourteen other individuals recombining between MA026a and D-Scar7 proved that the D locus was not localized between MA026a and D-Scar1 (Table 3). Two phenotyped individuals recombining between D-Scar0 and CPPCT040 reduced the interval containing the D locus: S5848-228 showed that the D locus was located upper than CPPCT040 while S6422-237 demonstrated that it was located below D-Scar0 (Table 3). Therefore, it can be concluded that the D locus is localized in a 0.4 cM interval between D-Scar0 and CPPCT040 (Fig. 4).

**Evaluation of the physical/genetic distance ratio around the D locus using a new BAC library**

The peach BAC library produced from F1-JF:21 DNA contained about 52,000 clones. Based on the analyses of a subset of clones, the average insert size was estimated at 90 kb, ranging from 50 to 130 kb. According to these preliminary results the covering of this BAC library was estimated at 15–16 × the peach haploid genome. D-Scar0 and D-Scar7 were used to screen the BAC library and four positive clones were found with D-Scar0 and 12 with D-Scar7. Three of the four positive clones with D-Scar0 were found to be also positive when screened with D-Scar7. The ratio between the genetic and physical distances was estimated using markers defined from BACend sequences of a positive clone common to both D-Scar0 and D-Scar7 (Table 4). A distance of 0.6 cM was estimated between the two CAPS markers F109-15-06 and R109-15-06 (Table 4) derived from the BACend sequences of one BAC containing an insert of 150 kb. This results in 1 cm corresponding to 250 kb in the region of the D locus.

**Discussion**

We describe in this paper major steps towards the cloning of the gene(s) controlling fruit acidity in peach, by phenotypic, genetic and physical characterization of the D locus.

The low polymorphism observed between 'Ferjalou Jalou sia™' and 'Fantasia' using AFLP markers was previously reported using RFLP and SSR markers [13] and was likely the consequence of the very low genetic distance between these two parental varieties [13]. Polymorphic markers would have been considerably increased by deep sequencing methods of the AFLPs [40]. Using the classical AFLP method in combination with bulk segregant analysis, 14 AFLP markers located within the 10 cM region harbouring the D locus were identified and no marker was mapped to another linkage group. These results confirmed that acidity trait in peach is not complex and should be controlled by a major gene.
Sequence analysis of the AFLP markers selected for conversion into SCAR revealed, at the vicinity of the D locus, an SNPs frequency 3.6 fold lower than the one reported in the 'Texas' × 'Earlygold' (TxE) reference Prunus map derived from an almond × peach interspecific cross (Illa E., personal communication).

To more accurately position the D locus, it was necessary to identify individuals of the extended population that had recombination events tightly linked to the D locus. The strategy followed in the present work involving two successive steps (firstly the genotyping of 1,718 individuals with the D locus-flanking markers and secondly the analysis of the recombinant individuals with additional tightly linked markers) reduced considerably the number of individuals that needed to be genotyped and phenotypically characterized. Fruit acidity is usually evaluated by pH or TA measurements. In this work, the use of both pH and TA was essential for the characterization of fruit acidity. In addition, the definition of thresholds based on the analyses of individuals without recombination event in the MA026a-BPPCT041 interval allowed a precise characterization of the phenotype. Thus, the phenotyping of the recombinant individuals with the pH and TA threshold strategy prevented misclassification of intermediate individuals that can be either homozygous (dd) or heterozygous (Dd) for the D locus.

The development of tightly linked markers and the phenotyping of recombinant individuals allowed the precise localization of the D locus. As fruit acidity is a major selection criterion, the D-linked markers could be used for marker assisted selection which would allow early selection of trees with the desirable character.

To estimate the relationship between the genetic and the physical distance at the vicinity of the D locus we anchored a BAC clone to the genetic map. Based on this analysis the ratio was estimated to 250 kb/cM. At the peach evergrowing (evg) locus the ratio was estimated to 10 to 35 kb/cM [41]. These ratios are smaller than the estimated average ratio on the TxE Prunus reference map [42] which is 553 kb/cM according to the genome size [2]. This is not surprising, as the physical/genetic distance ratio is known to vary along chromosomes [43-45]. The identification of the physical/genetic distance ratio in the vicinity of the D locus was important for estimating the number of walks needed for cloning the D locus. The D locus was localized in a 0.4 cM interval corresponding to a physical distance of 100 kb. Thus, one or two walks with BAC clones with an insert size of 90 kb should be sufficient to identify a BAC clone harbouring the D locus.

Sequenced BAC clones in peach [41,46], plum, apricot [46] and pear [47] revealed a gene density of 14 to 36 genes per 100 kb genomic sequence. Thus, in the 100 kb D locus region 14 to 36 candidate genes are expected. To identify the D gene(s) among these candidate genes, the aim will be to map accurately the recombination events relative to the predicted genes. To facilitate this analysis, the PP|FH BAC library was constructed from the F1-JF:21 hybrid between 'Ferjalou Jalousia®' and 'Fantasia' to identify one BAC clone for each allele. The two orthologous BAC clones will be sequenced and annotated and genetically dissected. In further analyses, the natural variability of the candidate genes will be explored within a peach germplasm collection to associate the haplotype to the phenotypic variation. Functional studies such as reverse genetics experiments should then provide further evidence for or against their involvement in fruit acidity.
Table 3: Genotypes and phenotypes of F2 recombinant individuals for nine markers framing the D locus

| Individual       | P1 | MA026a | D-Scar2 | D-Scar1 | D-Scar7 | D-Scar0 | CPPCT040 | D-Scar3 | D-Scar6 | BPPCT041 | No of F2 * |
|------------------|----|--------|---------|---------|---------|---------|----------|---------|---------|----------|-----------|
| SB220-1186       | [D] | H      | H       | H       | H       | H       | H        | F       | H       | F        | 15        |
| SB220-1321       | [d] | F      | F       | F       | F       | F       | F        | F       | F       | H        | 3         |
| SB48-350         | [D] | H      | H       | H       | H       | H       | H        | F       | F       | F        | 11        |
| SB220-1090       | [d] | F      | F       | F       | F       | F       | F        | F       | F       | H        | 3         |
| SB48-332         | [D] | H      | H       | H       | H       | H       | H        | F       | F       | F        | 2         |
| S642-022         | [d] | F      | F       | F       | F       | F       | F        | F       | F       | H        | 2         |
| S642-228         | [D] | H      | H       | H       | H       | H       | H        | F       | F       | F        | 1         |
| S642-237         | [D] | F      | F       | F       | F       | F       | H        | H       | H       | F        | 1         |
| SB220-2037       | [d] | F      | H       | F       | F       | F       | F        | F       | F       | F        | 1         |
| SB220-1188       | [D] | F      | F       | F       | H       | H       | H        | H       | H       | H        | 7         |
| S642-020         | [d] | H      | H       | F       | F       | F       | F        | F       | F       | F        | 2         |
| S548-147         | [D] | F      | F       | H       | H       | H       | H        | H       | H       | H        | 1         |
| S642-452         | [d] | H      | F       | F       | F       | F       | F        | F       | F       | F        | 1         |
| SB220-1045       | [D] | F      | H       | H       | H       | H       | H        | H       | H       | H        | 2         |

1 P: Phenotype, [d]: normally acid fruit, [D]: low acid fruit, 2 F: homozygous for the 'Fantasia' allele (dd), H: heterozygous highlighted in bold (Dd), * Number of F2 individuals having the same phenotype and genotype

The complete sequencing of the BAC clones will provide candidate genes for the D locus. These candidate genes may be structural genes implicated in metabolism or transport in agreement with our existing knowledge of fruit physiology or genes with novel structural or regulatory functions. Sequencing data will also provide information about Prunus genome organization in this particular region, which may be compared to homologous region in other Rosaceae species and even other fruit species. Microsystems analysis across Rosaceae species will provide insight into gene order, orientation and structural rearrangements of this particular region and through comparative genomics, may contribute to improve our knowledge on evolutionary and diversification processes among this family as demonstrated for Oryza [48].

Conclusion

In conclusion, the present work describes, for the first time, the fine mapping of a locus involved in a fruit quality trait on perennial plants via the chromosome landing approach. The development of tightly linked markers and the phenotyping of recombinant individuals allowed the precise localization of the D locus in a 0.4 cM interval corresponding to 100 kb. Using the constructed PPJFH BAC library with a mean insert size of 90 kb, one or two walks should be sufficient to identify a BAC clone harbouring the D locus. To our knowledge, only few fine genetic maps were realized using a large number of trees and only for resistance genes [49,50]. One of the major limitations for this strategy is the generation of a large population requiring an extended orchard maintained over several years. Our mapping population of 2,086 plants that segregates for many agronomic traits as well as the PPJFH BAC library will permit the genetic dissection of, not only the D locus, but also other traits such as Af (aborting fruit), S (flat/round fruit), G (peach/nectarine) and Ps (pollen sterility). This mapping population could be also exploited in any future genome sequencing project in peach where anchoring sequences or BAC contigs to the genetic map is a crucial step.

Methods

Plant material

The genetic linkage map was based on the segregation analyses of a peach F2 progeny. This progeny includes 208 individuals obtained from the selfing of a single F1 hybrid (F1-JF:21) issued from a cross between 'Ferjalou Jalousia®' a low-acid fruit variety and 'Fantasia' a normally-acid fruit variety. This population segregates for six Mendelian traits (low-acid/normally-acid fruit D, peach/nectarine G, flat/round fruit S, clingstone/freestone F, pollen sterility Ps, aborting fruit Af) [13] and for several characters involved in fruit quality as soluble sugar and organic acid concentrations [17]. Among the 208 individuals, 151 produced fruit at maturity while 57 produced flat fruit that fell after two months of growth and were not used in this study (Table 5).

For the fine mapping of the D locus, a total of 1,878 F2 additional individuals were obtained from the selfing of seven different F1 genotypes (Table 5). Three F1 individuals were issued from the cross between 'Ferjalou Jalousia®' and 'Fantasia' (F1-JF:21, F1-JF:28, F1-JF:104), two from the reverse cross (F1-FJ:47, F1-FJ:49), one from a cross between 'Fantasia' and 'Fercopale Platina®' (F1-PF:10) and one from the reverse cross (F1-PF:77). 'Ferjalou Jalousia®' is
homozygous for the dominant allele (DD), 'Fantasia' is homozygous for the recessive one (dd) and the seven F1 hybrids are heterozygous (Dd) for the D locus. 'Ferjalou Jalousia®' dominant allele is derived from 'Kiang-Si' that originated from China. 'Fercopale Platina®' and 'Ferjalou Jalousia®' shared the same common grandparents ('Kiang-Si' and 'Independence') (Fig. 6), and both had the same dominant allele for the D locus and produced flat peaches. According to a marker assisted selection for the Af gene that segregated in the 1,878 F2 individuals, 1,510 individuals were identified to produce fruit at maturity and were therefore genotyped. Among them, 1,084 individuals were planted in 2005 and 426 in 2006. The fine map was based on the genotyping of a total of 1,718 F2 individuals including the mapping population of 208 individuals and the 1,510 F2 additional individuals that should produce fruit at maturity (Table 5).

Fruit acidity phenotyping
The 151 F2 individuals of the mapping population producing fruit and recombinant individuals among the F2 progenies were phenotyped in 2007 and 2008. Two harvests separated by four days were performed for each individual. For each harvest, six fruits/individual were collected at maturity stage. TA and pH analyses were measured on fruit juice by using an equal volume of juice from each fruit as described previously [14].

To avoid any misclassification of recombinant individuals, we decided to rely on the analyses of homozygous and heterozygous individuals and to define thresholds in order to distinguish individuals with low-acid fruit from those with normally-acid fruit. Individuals, without recombination event, were selected on their genotype in the MA026a-BPPCT041 interval. Student's t-test was used to compare pH and TA mean values between homozygous and heterozygous individuals.

DNA extraction
Genomic DNA was extracted from young expanded terminal leaves. Fifteen milligrams of fresh weight were collected for each tree in 96 collection microtubes of 1.2 ml containing a tungsten carbide bead (3 mm diameter). They were ground in liquid nitrogen by using a Mixer Mill MM 300 (Retsch, Haan, Germany) for 1 min and 30 s and genomic DNA was extracted according to the method previously described [51].

BSA-AFLP
For Amplified Fragment Length Polymorphism (AFLP) assay combined with BSA, two low-acid (D/D or D/d) DNA bulks (BD1, BD2) and two normally-acid (d/d) DNA bulks (Bd1, Bd2) were used to identify putative markers linked to the D locus [52]. Individuals were selected among the 208 F2 used for the genetic linkage map according to juice pH and TA values measured in 2002 and 2003. Equal amounts of DNA from eleven individuals from the JxF F2 mapping population were pooled to construct each bulk.

The AFLP technique was performed following the protocol developed by [53] with some modifications. Genomic DNA (250 ng) was digested with two restriction enzymes PstI and MseI in a volume of 17.5 μl. The first PCR amplification was performed with primers having no selective nucleotide and then the second PCR amplification was carried out with primers having two selective nucleotides for PstI and three for MseI. PCR products were mixed with an equal volume of loading buffer (95% formamide, 30% deionized formamide, 0.05% bromophenol blue, 0.05% xylene cyanol, 0.5 M ammonium acetate) and loaded onto a 2% agarose gel. The gel was stained with ethidium bromide and photographed using a Gel Doc 2000 Imaging System (Bio-Rad, Hercules, CA). The RFLP bands were analyzed by using the BioNumerics software package (Applied Maths, Sint-Martens-Latem, Belgium). The RFLP data were used to construct an AFLP dendrogram using a UPGMA algorithm. The results showed that the AFLP markers were clustered into three main groups, one containing the normal acidity cultivars, one containing the low-acid cultivars and one containing the recombinant individuals. The main markers were used to design a reaction set for a genetic linkage map.
0.05% xylene cyanol, 0.05% bromophenol blue, 10 mM EDTA, pH 8.0). The mixture was heated for 5 min at 95°C, and then quickly cooled on ice. Each sample mixture was loaded on a 4.5% denaturing polyacrylamide gel and visualized by the silver staining system as described by [54]. Sixteen PsI+2 primers and 64 MsI+3 primers were tested consisting in a total of 1,024 primer combinations. Markers derived from PsI+2/MsI+3 primer combinations were named as pXX-YYY (X for the selective PsI nucleotides and Y for the selective MsI nucleotides). Subsequently, polymorphic AFLP primer combinations between bulks were used to screen the 208 F2 individuals of the mapping population.

Conversion of AFLP markers into SCAR and CAPS markers

AFLP markers linked to the D locus were selected for conversion into PCR markers for further easy use in large-scale screening of the 1,718 individuals and BAC library. After silver staining, marker fragments of the parents and of two F2 individuals were picked with a tip on the dried polyacrylamide gel [55] and dissolved in 15 μl deionized water. PCR amplifications were performed using 1 μl dilution with the same conditions as the selective PCR for AFLP reaction but with primers without selective nucleotides. The products were separated on a 2 to 3% agarose gel or 4.5% polyacrylamide denaturing gel.

Segregation analysis and map construction

Each polymorphic marker was tested by a chi-square for goodness of fit to the segregation ratios 1:2:1 expected for codominant markers and 3:1 expected for dominant markers in a F2 population. The linkage map was constructed using the MAPMAKER/EXP V3.0 software [56]. Markers were first divided into linkage groups using a crit-

Table 5: Origin of F2 individuals used for map construction and fine mapping of the D locus

| Cross | F1 name  | F2 name  | Number of F2 individuals |
|-------|----------|----------|--------------------------|
|       |           | S8220    | 208                      |
| F2 used for the construction of the genetic linkage map | F1-JF:21 | S8220 | 418 |
| 'Ferjalou Jalousia® × Fantasia' Producing fruits | F1-JF:28 | S6184 | 113 |
|       | F1-JF:104 | S7133 | 182 |
| 'Ferjalou Jalousia® × Fantasia' 'Fantasia' × Fantasia | F1-JF:47 | S6422 | 451 |
|       | F1-JF:49 | S6421 | 106 |
| 'Fantasia' × 'Ferjalou Jalousia®' | F1-FP:10 | S5848 | 405 |
| 'Ferjalou Jalousia® × Fantasia' 'Fantasia' × 'Fercopale Platina®' | F1-PF:77 | S6361 | 203 |
| Total of the additional F2 individuals | | | 1,878 |
| Additional F2 that will have fruits according to MAS for the Afgene | | | 1,510 |
| Total of the F2 individuals used for fine mapping | | | 1,718 |

Figure 6

Origin of ‘Ferjalou Jalousia®’, ‘Fantasia’ and ‘Fercopale Platina®’ peach varieties. The phenotype for the D locus is indicated for each variety. Varieties with [D] phenotype produce low-acid fruit and varieties with [d] phenotype produce normally-acid fruit.
ical LOD score threshold of 5. The Kosambi function was used to convert recombination units into genetic distances.

**Fine mapping**

The 1,510 additional individuals that would have fruit at maturity were used to complete the mapping population to a total of 1,718 individuals segregating for the D locus. These individuals were screened for five markers, two SSR markers and three new SCAR markers, spanning a large region around the D locus: MA026a and BPPCT041, previously mapped on JxF linkage map [13] and three AFLP markers transformed into SCAR markers. Recombinant individuals detected in this region were genotyped with three other AFLP markers transformed into SCAR markers and CPPCT040 a SSR marker mapped on the top of the linkage group 5 of 'Texas' × 'Earlygold' linkage map [26]. The phenotype of the recombinant individuals compared to the recombination point enabled the localization of the D locus. Among the 308 recombinant individuals detected, 149 individuals producing fruit in 2007 and 2008 were phenotyped for fruit pH and TA.

**Bacterial Artificial Chromosome (BAC) library construction**

The *Prunus persica* PPJFH BAC DNA library was realized at URGV (INRA, Evry) and constructed as described previously [57]. Nuclei were isolated from 32 g of young leaves frozen in liquid nitrogen from F_1 hybrid DNA heterogeneous for all the mendelian characters segregating in the JxF cross. Restriction fragments were subjected to a double size selection in a CHEF-DRIII apparatus (Bio-Rad) by pulse field gel-electrophoresis (PFGE). The DNA from the agarose slices was electrophoresed and cloned into the pIndi pulse field gel-electrophoresis (PFGE). The DNA from the size selection in a CHEF-DRIII apparatus (Bio-Rad) by JxF cross. Restriction fragments were subjected to a double zygous for all the mendelian characters segregating in the 2008 were phenotyped for fruit pH and TA.

To associate a genetic distance to the physical distance obtained with the PFGE, primers were designed using Primer3 software (version V.0.4.0) based on two BACend sequences of a chosen positive BAC clone for two markers (D-Scar0 and D-Scar7). The primers of two markers were tested for PCR amplification on 'Ferjalou Jalouzia®' and 'Fantasia' to identify polymorphism between the parents. Reaction mixtures and PCR conditions were done as described for SCAR markers and the amplified fragments were then sequenced. Obtained markers were used to genotype only individuals recombining between D-Scar1 and CPPCT040 framing D-Scar0 and D-Scar7.

**Authors’ contributions**

KB carried out the molecular genetic studies, the sequence alignment and drafted the manuscript. KB and AB conceived and designed the experiments for the BAC library. GC participated to AFLP mapping. CT carried out the BAC library pooling and extraction. KB, GC, AM and ED performed the phenotypic analysis. ED and GC participated in the genetic studies. AB and ED conceived the study and participated in its design. AB, AM and ED helped to draft the manuscript. All authors read and approved the final manuscript.

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