

Identification and validation of a QTL for fruit firmness on linkage group 4 in three sweet cherry populations

Lichun Cai, José Quero-Garcia, Teresa Barreneche, Elisabeth Dirlewanger, Christopher Saski, Amy Iezzoni

► To cite this version:

Lichun Cai, José Quero-Garcia, Teresa Barreneche, Elisabeth Dirlewanger, Christopher Saski, et al.. Identification and validation of a QTL for fruit firmness on linkage group 4 in three sweet cherry populations. 9th International Rosaceae Genomics Conference, Jun 2018, Nanjing, China. hal-03358885

HAL Id: hal-03358885 https://hal.inrae.fr/hal-03358885

Submitted on 29 Sep 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

ROSBREED COMBINING DISEASE RESISTANCE WITH HORTICULTURAL QUALITY IN NEW ROSACEOUS CULTIVARS

Identification and validation of a QTL for fruit firmness on linkage group 4 in three sweet cherry populations

Lichun Cai¹, José Quero-Garcia², Teresa Barreneche², Elisabeth Dirlewanger², Christopher Saski³, Amy Iezzoni¹ ¹Department of Horticulture, Michigan State University, East Lansing, MI, 48824, USA; ²UMR 1332 BFP, INRA, Université de Bordeaux, F-33140 Villenave d'Ornon, France; ³Department of Plant and Environmental Sciences, Clemson University, Clemson, SC, 29634, USA

Introduction

Fruit firmness is an important market driven trait in sweet cherry (*Prunus avium* L.) as consumers prefer firm sweet cherries (3). The firmness exhibited by modern sweet cherry varieties is a derived trait as wild cherry has small soft fruit. These increases in fruit size and firmness are the two major traits associated with domestication. However, little is known about the genetic control of fruit firmness in sweet cherry. Therefore, the objectives of this study were to identify and characterize loci that control fruit firmness in sweet cherry from three different germplasm populations, one of which contain representatives from wild cherry.

Materials and Methods

- Three sweet cherry populations were evaluated in this study: Pop1- an INRA biparental F₁ population from the cross of 'Fercer' × 'X' (X=unknown parent), Pop2 an INRA diversity population, and Pop3 - RosBREED's pedigreed populations.
- Fruit firmness was evaluated using Durofel® and BioWorks' FirmTech 2 texture analyzer for INRA and RosBREED populations, respectively. ✤ All the individuals were genotyped with RosBREED Illumina Infinium® cherry SNP array (2).
- ☆ MultiQTL (MultiQTL Ltd, Haifa, Israel) and FlexQTL[™] (1) softwares were used for QTL mapping in the F₁ and RosBREED populations, respectively. A general linear model was employed to test the marker-trait association in the diversity population.

Fig. 1. Frequency distributions of fruit firmness for the three sweet cherry populations: (a) Pop1 (n=67); (**b**) Pop2 (n=193); (**c**) Pop3 (n=528).



Results

Fig. 2. The major QTL identified in the 'Fercer' \times 'X' F₁ population mapped to the 'X' linkage group 4. Explained percentage of variance is given for each QTL. This previously unidentified QTL that was named *qP-FF4.1*.

Fig. 3. Association of chromosome 4 SNP markers with fruit firmness in the INRA diversity population. (a) P values for all the SNP markers on chromosome 4. (b) Phenotypic means based on the genotype of two SNP markers that fall within the *qP-FF4.1* interval, the SNP marker ss4905522928, and (c) the SNP marker ss490552906. Phenotypic means that were significantly different (P < 0.05) are identified by different letters.

MICHIGAN STATE

UNIVERSITY





Fig. 4. *qP-FF4.1* was identified in the RosBREED pedigreed population. The QTL positions along genome estimated by FlexQTL[™] the are illustrated by trace plots (top) and posterior probabilities (bottom).



Fig. 5. Haplotypes for the fruit firmness QTL, qP-FF4.1. Haplotypes only identified in wild sweet cherry are marked as blue. The haplotypes were deduced to be associated with soft or firm fruit based on diplotype analysis

Marker	SNP	ss490552883	ss490559054	ss490552906	ss490552912	ss490552928
	Physical postion (bp)	10,241,247	10,414,884	10,880,163	11,044,975	11,472,398
	Genetic position (cM)	33.5	33.6	33.7	33.7	33.8
	H1 (firm)	В	В	А	В	В
	H2 (soft)	А	В	В	А	А

Fig. 6. Comparison of fruit firmness for progeny based on their diplotypes for qP-FF4.1 from (a) 'Fercer' × 'X' F_1 population; and (b) RosBREED pedigreed population. Only diplotypes represented by six or more individuals were included. Phenotypic means that were significantly different (P < 0.05) are



Conclusions and Future Work

A newly identified QTL for fruit firmness, named *qP-FF4.1*, was identified on LG4 in all three sweet cherry populations. The "soft fruit" alleles for *qP-FF4.1* contributed by the wild 'mazzard' cherries were dominant over the "firm fruit" alleles. These results advance our understanding of the genetic basis of fruit firmness and will help to enable the use of DNA informed breeding for this trait in sweet cherry breeding programs. Future work will include the identification of candidate genes within the QTL interval.

References

- Bink et al. 2014, Theor Appl Genet 127(5):1073-1090
- Peace et al. 2012, PLoS ONE 7(12):e48305 2.
- Yue et al. 2017, Agriculture and Resource 3.

Economics Review 46:103-122



Funding for RosBREED: Combining disease resistance with horticultural quality in new rosaceous cultivars is provided by the Specialty Crop Research Initiative Competitive Grant 2014-51181-22378 of the USDA's National Institute of Food and Agriculture.

We thank the INRA's Prunus Genetic Resources Center for preserving and managing the sweet cherry collections and the Fruit Tree Experimental Unit of INRA-Bordeaux for maintaining the trees.